Pathways from socioeconomic deprivation to bronchiolitis and subsequent childhood asthma

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A thesis submitted for the degree of Doctor of Philosophy

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DECLARATION

I, Kate Marie Lewis, confirm that the work presented in this thesis is my own. Where information has been derived from other sources, I confirm that this has been indicated in the thesis.

Date: ___________________________ Signature: ___________________________
ABSTRACT

**Introduction:** Bronchiolitis and childhood asthma are major causes of morbidity among children in the UK, yet there are no preventative or curative measures for most children that develop these conditions. A better understanding of the longitudinal pathways to these conditions is warranted to design effective prevention policies. Using a social determinants of health framework, I explored the pathways between socioeconomic position, bronchiolitis and childhood asthma.

**Methods:** I used national birth cohorts created from linked administrative datasets in my thesis. I used harmonic Poisson regression models to examine associations between socioeconomic deprivation and the seasonality of bronchiolitis admissions in England. I modelled typical trajectories of asthma/wheeze among children in Scotland using latent class growth analysis. Using causal inference methods, I estimated: the socioeconomic disparities in the risk of bronchiolitis admissions that would remain if maternal smoking during pregnancy were eliminated; and the socioeconomic disparities in the risk of chronic trajectories of asthma that would remain if bronchiolitis admissions were eliminated.

**Results:** The peak timing of bronchiolitis admissions varied marginally across England, with earlier peaks in areas with higher population densities. After accounting for seasonal patterns, the North of England had disproportionately higher rates of admissions and, nationwide, disparities followed a socioeconomic gradient. I estimated that eliminating maternal smoking would reduce 20% of socioeconomic disparities in the risk of bronchiolitis admission. I identified four asthma/wheeze trajectories in children: no/infrequent, early-transient, early-persistent and intermediate-onset. Eliminating bronchiolitis admissions could reduce up to 18% of the disparities in the risk of chronic asthma by age ten.

**Conclusions:** Intervening early on the most socioeconomically deprived populations should be central to policies aiming to reduce the incidence of bronchiolitis admissions and asthma. The contribution of other socioeconomically patterned risk factors, including pollution and housing conditions, should be investigated in future work.
IMPACT STATEMENT

The work in this thesis addresses two key child public health issues in the UK. Bronchiolitis, the leading cause of hospital admission in infancy, and asthma, the most common chronic respiratory condition in children. Other epidemiological studies have addressed these conditions (and the potential link between the two), but none with the systematic methods presented here. I focus on socioeconomic deprivation, adding to the evidence base showing a socioeconomic gradient in the incidence of bronchiolitis and paediatric asthma in the UK. Further, my work begins to untangle some of the pathways through which inequities in these conditions develop.

Drawing from the conceptual framework developed from the World Health Organisation's Commission of the Social Determinants of Health, I create a structure to study the complex layers of risk factors that influence the development of bronchiolitis and subsequent asthma. My work includes two studies that look at the mechanisms of the seasonality of bronchiolitis admissions using epidemiological methods. The seasonal pattern of bronchiolitis has long been recognised by paediatricians, but there is no research to date that has studied and quantified this effect in a UK setting. I then use causal inference methods to quantify the contribution of specific risk factors in the development of bronchiolitis and asthma. These findings can be used to enhance public health strategies towards the alleviation of inequity in bronchiolitis and asthma.

I provide an overview of causal mediation estimands recently proposed in the literature, critically assessing the potential application of each to health inequity research using observational data. As part of this, I apply a novel method of evidence synthesis to design causal diagrams to inform analysis. I show the complexities involved in the process of finding unbiased estimates of mediation effects, but also provide some workable solutions to this. This work provides a framework through which other researchers can apply these methods to health inequity research in the future. I have made the code for the statistical analyses in my published studies freely and publicly available on GitHub. This is important for transparency and reproducibility of research, and I will continue to do this for further papers published from my thesis. I have published three open-access journal articles directly based on work presented in this thesis and have presented my findings at several conferences.

I list specific outputs produced over the course of my PhD in Appendix 0, p.195.
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<tr>
<td>A&amp;E</td>
<td>Accident and emergency</td>
</tr>
<tr>
<td>aOR</td>
<td>Adjusted odds ratio</td>
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<tr>
<td>BIC</td>
<td>Bayesian information criterion</td>
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<tr>
<td>BLRT</td>
<td>Bootstrapped likelihood ratio test</td>
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<tr>
<td>BNFC</td>
<td>British National Formulary for Children</td>
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<tr>
<td>CCG</td>
<td>Clinical Commissioning Group</td>
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<tr>
<td>CDE</td>
<td>Controlled direct effect</td>
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<tr>
<td>CDM</td>
<td>Counterfactual disparity measure</td>
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<tr>
<td>CHI</td>
<td>Community Health Index</td>
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<tr>
<td>CI</td>
<td>Confidence interval</td>
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<tr>
<td>COPD</td>
<td>Chronic obstructive pulmonary disease</td>
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<td>CSDH</td>
<td>Commission on Social Determinants of Health</td>
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<tr>
<td>DAG</td>
<td>Directed acyclic graph</td>
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<td>eDRIS</td>
<td>Electronic Data Research and Innovation Service</td>
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<tr>
<td>ESC-DAG</td>
<td>Evidence synthesis for constructing directed acyclic graphs</td>
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<tr>
<td>GP</td>
<td>General practitioner</td>
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<td>GMM</td>
<td>Growth mixture modelling</td>
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<td>GWAS</td>
<td>Genome-wide association study</td>
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<td>HES APC</td>
<td>Hospital episode statistics admitted patient care</td>
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<td>HR</td>
<td>Hazard ratio</td>
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<td>IMD</td>
<td>Index of multiple deprivation</td>
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<td>IRR</td>
<td>Incidence rate ratio</td>
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<td>ISAAC</td>
<td>The International Study of Asthma and Allergies in Childhood</td>
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<tr>
<td>IPW</td>
<td>Inverse probability weighting</td>
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<td>LCGA</td>
<td>Latent class growth analysis</td>
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<td>LSOA</td>
<td>Lower super output area</td>
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<tr>
<td>MHCLG</td>
<td>Ministry of Housing, Communities and Local Government</td>
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<tr>
<td>MICE</td>
<td>Multiple imputation by chained equations</td>
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<tr>
<td>MSM</td>
<td>Marginal structural model</td>
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<td>NDE</td>
<td>Natural direct effect</td>
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<td>NHS</td>
<td>National Health Service</td>
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<td>National Institute for Health and Care Excellence</td>
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<td>NIE</td>
<td>Natural indirect effect</td>
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<td>NO₂</td>
<td>Nitrogen dioxide</td>
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<td>Full Form</td>
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<td>NRS</td>
<td>National Records of Scotland</td>
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<td>NS-SEC</td>
<td>National Statistics Socio-Economic Classification</td>
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<td>ONS</td>
<td>Office for National Statistics</td>
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<td>OR</td>
<td>Odds ratio</td>
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<td>PHE</td>
<td>Public Health England</td>
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<td>PHS</td>
<td>Public Health Scotland</td>
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<td>PICU</td>
<td>Paediatric intensive care unit</td>
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<td>PIS</td>
<td>Prescription Information System</td>
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<td>PM$_{2.5/10}$</td>
<td>Particulate matter (less than 2.5/10 micrometres in diameter)</td>
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<td>RIA</td>
<td>Randomised interventional analogues</td>
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<td>RRR</td>
<td>Relative risk ratio</td>
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<td>RSV</td>
<td>Respiratory syncytial virus</td>
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<td>SBR</td>
<td>Scottish Birth Records</td>
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<td>SGA</td>
<td>Small for gestational age</td>
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<td>SEP</td>
<td>Socioeconomic position</td>
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<tr>
<td>SIGN</td>
<td>Scottish Intercollegiate Guidelines Network</td>
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<td>SIMD</td>
<td>Scottish index of multiple deprivation</td>
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<td>SMR</td>
<td>Scottish Morbidity Record</td>
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Chapter 1. Background

Chapter overview

The overarching aim of Chapter 1 is to present an overview of the current literature and public health policies relevant to this thesis, focusing on the relationships between socioeconomic deprivation, bronchiolitis and the subsequent development of asthma. Firstly, I will present an overview of the epidemiology of bronchiolitis and paediatric asthma in the United Kingdom, and introduce how the two conditions may be linked. Where possible I will focus specifically on England and Scotland, the nations from which my study data originates. Secondly, I will define health inequity and socioeconomic position—key terms in my work—and explain the conceptual framework that underpins this research. Thirdly, I will present an overview of pre- and post-natal risk factors for bronchiolitis and asthma. Lastly, I will outline current policy and practice relevant to the prevention and treatment of bronchiolitis and childhood asthma in England and Scotland, with a particular focus on efforts to lessen socioeconomic inequities in these conditions.

1.1. Introduction

1.1.1. Bronchiolitis

Bronchiolitis is an acute lower respiratory tract illness caused by viral infections that affects young children, particularly infants.\(^1\) Diagnosis of bronchiolitis is based on respiratory signs and symptoms rather than a diagnostic test or universal scoring system.\(^2,3\) This infection typically begins with symptoms similar to the common cold, but progresses to a persistent cough, fast and shallow breathing, chest recession and crackles and/or wheezing within a few days.\(^1,3\) Mild cases of bronchiolitis can be managed at home, but a significant number of infants require supportive hospital care to aid feeding and respiration.\(^4,5\) In infants admitted to hospital in England with bronchiolitis, approximately 80% of cases are caused by respiratory syncytial virus (RSV),\(^6\) however, other respiratory viruses including human metapneumovirus (HMPV), rhinovirus, adenovirus, influenza and parainfluenza are also known to cause bronchiolitis.\(^1\) These viruses are mainly transmitted through direct and indirect contact (via contaminated intermediate objects) with an infected person and by airborne droplets, which are spread through coughing, sneezing and talking.\(^7\) There is early evidence to suggest that
RSV may also be spread by aerosolised (airborne) particles, which are smaller than droplets and have a longer range of transmission.\(^8\)

Whilst mortality due to this condition is uncommon in the United Kingdom (UK) (in 2019 there were 7 deaths in England and 2 deaths in Scotland among young children with bronchiolitis recorded as the underlying cause),\(^9,10\) its associated morbidity is substantial. In 2018/19, there were 48,744 hospital admissions in England with a primary diagnosis of bronchiolitis among infants (children <1 year old), making it the most common reason for hospital admission in the first year of life.\(^11\) Qualitative research from Canada and France highlights the emotional strain, disruption and anxiety bronchiolitis places on the families of infants hospitalised with this infection.\(^12\)–\(^14\) The rate of hospital admissions for this condition have been rising substantially over time,\(^4\) leading to warnings about needing to “stem the rising tide” of these admissions.\(^15\)

RSV-attributable bronchiolitis in children less than 2 years old is also estimated to contribute an more than 82,000 general practitioner visits annually in the UK.\(^16\) Reflecting the seasonal pattern of RSV, cases of bronchiolitis peak in early winter in the UK, contributing to pressures on the National Health Service (NHS) during a particularly stretched time of year.\(^1,4\)

1.1.2. Asthma

Asthma, a respiratory disease involving airway inflammation and hyper-responsiveness, is one of the most common chronic conditions among children in the UK.\(^17\) In 2019, the estimated prevalence of asthma (defined by self-/parent-reported doctor’s diagnosis of asthma and wheeze in the last year) among under 15’s in the UK was 9.6% (95% confidence interval (CI) 6.5 to 13.7).\(^18,19\) This compares to a Europe-wide average of 5.7% (95% CI 3.9 to 8.1) and a global average of 4.7% (95% CI 3.3 to 6.7). Symptoms of asthma can vary over time and with intensity, but commonly include wheezing, chest tightness, breathlessness and coughing.\(^20,21\) Exacerbations of symptoms, commonly known as an asthma attack, can lead to variable airflow obstruction and inflammation within the lungs, which may be fatal if left untreated.\(^17\) It is increasingly recognised that asthma is not a singular disease but an umbrella term for a heterogeneous group of conditions.\(^22,23\) Relatedly, there is no single gold standard definition or diagnostic test for asthma.\(^17\)

Different trajectories of childhood asthma, defined by the evolution over time in reported symptoms, have been distinguished based on longitudinal data.\(^24\)–\(^26\) This includes early-transient asthma (beginning before age 3 and subsiding after a few years), early-persistent asthma (beginning before age 3 and continuing into later childhood) and intermediate/late-onset asthma (beginning between 3 and 8 years). In an estimated 40-60% of cases, allergic sensitisation is also a characteristic of the disease; a category of asthma sometimes defined
as atopic asthma. Early asthma symptoms may be indicative of viral or transient wheeze caused by respiratory infections and, therefore, diagnosis of asthma is not recommended in children under the age of 5. Where symptoms are still present at age 5, the National Institute for Health and Care Excellence (NICE) recommends that clinicians use objective tests (e.g. spirometry) to then diagnose asthma in these children.

The direct costs for asthma across all NHS services has been estimated to be £1.1 billion, including 85,000 hospital admissions and 108,000 general practitioner (GP) consultations across all age groups per year in the UK. In 2018/19, there were 25,342 recorded hospital admissions with a primary diagnosis of asthma among under-18s in England. Asthma-related deaths in children are rare, with 15 deaths in England and 1 death in Scotland among 0-19 year olds with asthma recorded as the underlying cause in 2019. However, these deaths are almost entirely, if not completely, avoidable. Of the 28 deaths in people less than 20 years old assessed in the 2014 National Review of Asthma Deaths, 26 (or 93%) were ascertained to be "potentially avoidable". Further, data published by the Global Health Data Exchange shows that, in 2017, the UK had the second highest rate of death from asthma amongst 15-19 year olds in a comparison with 18 other high income countries. Childhood asthma is associated with other adverse outcomes, including an increased risk of anxiety and depression, more absentee days from school and poorer educational outcomes than peers, and diminished self-rated quality of life.

1.1.3. Why focus on these two conditions?

Despite being the cause of substantial morbidity and cost to the NHS, there are currently no preventative or curative measures for the majority of children that develop either bronchiolitis or asthma.

Under UK guidelines, infants at high risk of RSV-related hospital admissions, including preterm infants with chronic heart or lung conditions that are less than six months of age at the beginning of the RSV season, are recommended to receive palivizumab, a humanised monoclonal antibody, prophylactically during peak months of RSV circulation. Costing between £3000 and £5000 to treat each child with a full course, it is not deemed cost-effective to administer this preventative treatment to the wider infant population (who, on the majority, will not develop severe symptoms from RSV infection). There is also no vaccination currently available against RSV, the dominant pathogen causing bronchiolitis, although several candidates are currently being assessed in clinical trials. Given that approximately 80% of children admitted to hospital in England with bronchiolitis are reported as otherwise healthy,
this means that there is no preventative healthcare interventions for the majority of infants at risk of bronchiolitis.44,45

In the last decade, previous gains in improving key outcomes for asthma globally, including reducing mortality and hospital admission rates, have reportedly stalled.46,47 Asthma care focuses on managing symptoms through the use of medicines that prevent and relieve symptoms, such as inhaled corticosteroids.20 Poor management of symptoms, particularly around the use of preventative medicine, is referenced at the outset of the Scottish Intercollegiate Guidelines Network (SIGN) guidelines on the management of asthma as the driver of “much of this morbidity”.48 This document highlights the failure of primary prevention methods, including allergen avoidance, modified milk formulae, weaning and nutritional supplementation, to offer clear beneficial outcomes on asthma outcomes. Breastfeeding, childhood immunisations and reduced smoking exposure are recommended for all children (and weight loss interventions for overweight and obese children), but as factors that “promote general health” rather than specifically in relation to preventing asthma. Trials of pharmacological prevention using antihistamines have been undertaken but here is little evidence to show that these stop the long-term development of asthma so far.20

A better understanding of the risk factors for these conditions is therefore imperative to plan how to best intervene early and reduce the incidence and population impact of these conditions in children. Prominently, severe symptoms of RSV infection have been consistently associated with an increased risk of wheeze and asthma in later childhood across the research literature.49–53 For example, infants born in English NHS hospital in 2007/08 were 4.4 times more likely to have a hospital admission for asthma between ages 1 and 5 if they had one or more admissions for bronchiolitis in infancy, compared to those with no admission for bronchiolitis (adjusted hazard ratio (HR) 4.35, 95% CI 4.00 to 4.73).54 There is a debate to whether this association represents a true causal mechanism (i.e. early infection impairs pulmonary function, thereby directly influencing the development of asthma) or it is due to other shared influences, such as environmental exposures or a familial disposition to respiratory ill health.49,51,55,56 Notable recent evidence from the MAKI randomised controlled trial in the Netherlands has shown the effectiveness of palivizumab prophylaxis, given to otherwise healthy preterm infants, in reducing wheeze up to age 3 but not at age 6.57,58 Or, to put this another way, RSV infection as an important mechanism for early, but not later, wheeze.57,58 These results further open up the question of alternative—and potentially modifiable—factors that may jointly influence the development of both conditions.55
My PhD will focus specifically on the common socioeconomic determinants of bronchiolitis and asthma to elucidate the pathway to and between the two conditions, and the potential for interventions, further.

### 1.2. Health inequities and socioeconomic position

#### 1.2.1. Key definitions

Health inequity and socioeconomic position (SEP) are concepts that I will refer to often throughout this thesis. By inequities in health, I mean the disparities in health outcomes (or in the opportunity to be healthy) across strata of society that are created by systematic social, political and economic conditions. I use the term health inequity as opposed to inequality to emphasise the avoidable and unjust nature of these differences in circumstance and health outcomes. Within my PhD different social strata will be characterised by SEP, defined by Galobardes and colleagues as “the social and economic factors that influence what positions individuals or groups hold within the structure of a society”. Socioeconomic deprivation will be used to refer to the proportion of the population at the most deprived end of SEP.

#### 1.2.2. Measuring SEP

Although commonly employed in health research, there is no agreed upon definition of such a multidimensional concept such as SEP, which may encompass education, occupation, income, ownership of goods, social prestige and other factors. Therefore, there is no standard method of measuring SEP across health inequity or epidemiological studies. As argued by Bartley, the definition of SEP used should be guided by research hypotheses. For purposefully designed studies on health inequity, careful consideration of how to measure SEP and related concepts may be achieved. Unfortunately, as in many cases, the measurement of SEP in my PhD, is driven by the available data. However, by discussing the available indicator at the outset of my thesis, I hope to be transparent about the conclusions about SEP that I can make in my research based on my employed measure.

**My measure of SEP: the Index of Multiple Deprivation**

The Index of Multiple Deprivation (IMD) is a relative measure of deprivation across small areas, constructed by assigning each area a value calculated from weighted scores across seven domains of deprivation: income, employment, health, education, access to services, living environment and crime. England and Scotland have their own versions of IMD, which differ in respect to the coverage of each small area (approximately 1500 residents in England
Compared to 760 in Scotland) and the precise components of the seven domains.\textsuperscript{63,64} IMD is an official statistic, constructed by the Governments of England and Scotland for a variety of purposes, including: to shape policy and strategy; identify priority areas to target resources; and monitor inequalities across small areas.\textsuperscript{65} The indices are updated every 4 to 5 years to reflect changes in the underlying data, with the same methodology and domains used.\textsuperscript{65} There are two other notable area-level measures of material deprivation that cover the whole of the UK; the Carstairs Index and Townsend Deprivation Index.\textsuperscript{66} These indices are both derived from components of census data including car ownership, unemployment, overcrowding, low social class (Carstairs Index only) and home ownership (Townsend Index only).

I will now outline the strengths and limitations of IMD with reference to the two goals of measuring inequity in my work.\textsuperscript{67}

\textit{Goal one: to monitor and describe the social distribution of disease}

Monitoring the social determinants linked to health outcomes is an essential step to changing the course of health inequities. IMD is an apt measure for monitoring as it provides a consistent and nationally applicable measure of inequity. IMD has been used extensively in health inequity research, including in studies of bronchiolitis and asthma,\textsuperscript{4,66,69} which means that my results can be compared with previous and future work. IMD captures a wide range of factors that contribute to inequity, which is arguably more suited to the similarly comprehensive concept of SEP than a non-composite measure. In addition, IMD is an area-level measure, which means it can characterise some living conditions that individual or household measures cannot, such as ambient air pollution.\textsuperscript{70} On the other hand, use of this ecological measure as a proxy for an individual level indicator can, in some circumstances, overestimate the true individual-level effect of SEP.\textsuperscript{67} Notably, as IMD was constructed to measure specific aspects of deprivation rather than affluence, the indices are less able to discriminate between areas in the less deprived end of the distribution.\textsuperscript{63} A criticism levelled at the IMD’s identification of the most deprived areas is its heavy reliance on benefit-recipient data, which means it undercounts “entitled nonrecipients”\textsuperscript{.71}

\textit{Goal two: to identify interventions to weaken the relationship between socioeconomic deprivation and disease}

There are two feasible ways to go about decreasing socioeconomic inequities in health outcomes. The first is by directly intervening on SEP and decreasing overall socioeconomic inequity across society. This is an ambitious but not impossible goal, with Heyman writing that this “major social change requires sustained public support and political will”.\textsuperscript{72} p.369 A universal
basic income and formal (high quality) early years education and childcare are widely promoted ways to reduce inequities.\textsuperscript{73,74} The 2010 Marmot Review advocates the universal application of actions such as these, but at a scale and intensity that varies proportionately across the social gradient, a method coined as proportionate universalism.\textsuperscript{75} Other measures of SEP, such as income or education, are arguably more suited to measuring this type of policy change due to their specificity. However, a composite measure of SEP may be a better indicator of the broader societal change described by Heyman.\textsuperscript{72} IMD in particular, though, may not be the best measure of changes within areas because it is a relative measure of deprivation and is not suited to observing absolute change in deprivation over time.\textsuperscript{63}

The second approach to reducing health inequities is to intervene on risk factors that sit on the pathway between SEP and disease (for example, by improving housing conditions or reducing rates of smoking). This, arguably, more pragmatic and achievable approach is what my thesis focuses on. To achieve the goal of identifying interventions to reduce inequities, we need to first quantify the extent to which risk factors contribute to health inequities. As a nationally applicable measure of SEP, IMD is a helpful way of categorising the population by levels of socioeconomic deprivation to further investigate these underlying risk factors. Indeed, IMD was designed as a tool for local policy makers and communities to identify the most deprived areas for “the effective targeting of resources”.\textsuperscript{64} On the other hand, some of the components that are used to create IMD may include indicators of the very risk factors that we wish to investigate. For example, the living environment domain includes measures of the quality of housing and air pollution. To avoid the potential of ‘mathematical coupling’ (whereby two variables correlate if one shares part of the other), some researchers choose to remove the health domain from IMD when conducting health inequity research.\textsuperscript{70,76}

### 1.3. Conceptual framework

The pathways between SEP and bronchiolitis, and later asthma, are highly complex. To develop specific, yet policy relevant, research questions that can be answered with the available data, I will draw from the multifaceted conceptual framework developed for the Commission on Social Determinants of Health (CSDH) by the World Health Organisation (WHO).\textsuperscript{60} A key strength of the CSDH framework lies in the differentiation between levels of causation. This framework purports that the socioeconomic and political context of a society creates socioeconomic strata by factors such as income, education, occupation, gender and/or ethnicity. In turn, these structural dimensions operate through intermediary factors, which broadly include material and psychosocial circumstance and behavioural factors, to shape health outcomes.
Figure 1.1, which I have adapted from the CSDH, displays the guiding framework of this PhD in reference to inequity in bronchiolitis and asthma outcomes. Alongside SEP, which is the structural determinant of inequity central to my thesis, I include neighbourhood, built environment, and geographical location and climate. These environmental dimensions are strongly associated with socioeconomic factors, but are also independently associated with risk factors important to respiratory disease, such as pollutants and viruses. In addition, although not explicitly displayed in the framework, the life course perspective is an essential component of the social determinants of health. This perspective helps to frame the temporal element to my project; namely, that exposures earlier in life contribute to respiratory disease outcomes months or years later in life. The effects of risk factors may accumulate over life and/or specific insults may happen during a critical or sensitive period, leading to long-lasting effects on later disease outcomes. For example, children with asthma by the age of seven have been shown to have lung function deficits at just one month of age, emphasising the early origins of at least some asthma phenotypes.

In the next sections, I will review the evidence for each component of my framework in turn.

1.3.1. Literature search strategy

The information presented in this narrative review of the literature was initially selected by searching electronic databases (Medline, Embase, Maternity and Infant care database and
PubMed) and national statistics with keywords relevant to each section. For example, bronchiolitis and ethnicity, or asthma and smoking. I applied a snowball method (pursuing references of references) to these papers to find the literature most relevant to describing inequities in bronchiolitis infections and asthma in the UK. To ensure that I also included the most recent research available in the field, I subsequently added to the review using the papers identified using the search criteria outlined in Table 1.1.

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<th>Table 1.1. Literature search terms for two overarching research questions</th>
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1.4. **Structural determinants**

In this section, I will present evidence on the association between structural determinants of bronchiolitis and asthma in a UK context. Whilst the primary focus of my PhD is on SEP, I will also introduce evidence on the relationship between ethnicity and immigration status—other structural determinants that interact closely with SEP to create inequities—and these health outcomes in this section. Geographical location, the built environment and neighbourhood are included here as additional factors that interact with SEP, as well as work independently, to influence the development of bronchiolitis and asthma.

### 1.4.1. SEP

Evidence from several studies of administrative health data show a socioeconomic gradient in the incidence of severe bronchiolitis (i.e. cases that require hospitalisation), a pattern that has persisted over many years. Using data from the Oxford Record Linkage Study, Green et al. looked at hospital admissions among children born between 1970 and 1989. After adjustment for other risk factors, they found a 2.34 greater odds of admission to hospital for bronchiolitis among infants born into the lowest fifth of SEP (measured using the Registrar General's occupational classification, based on the head of the household), compared to the highest fifth of SEP (odds ratio (OR) 2.34, 95% CI 1.65 to 3.30). Among births in England between 2003 and 2012, infants with a birth address in the highest IMD decile had a 1.94 greater incidence of hospitalisation for bronchiolitis (sex and year-adjusted incidence rate ratio (IRR) 1.94, 95% CI 1.73 to 2.19) and those in the 50%-89% of IMD a 1.48 greater incidence rate (sex and year-adjusted IRR 1.48, 95% CI 1.07 to 2.06), compared to infants in the lowest IMD decile. A similar pattern but at a lower risk was observed among infants born in Scotland over the same period (sex and year-adjusted IRR 1.28, 95% CI 1.16 to 1.42, highest compared to the lowest Carstairs index decile). The Carstairs index is constructed at the postcode sector level, covering a population on average more than 3 times the size of the English IMD, which likely explains the lower risk found in Scotland compared to England.

A study of all 29 Paediatric Intensive Care Units (PICUs) in England and Wales between 2004 and 2007 also found that admissions for acute respiratory failure and bronchiolitis were more likely among infants from the most deprived fifth of the community (measured using the Townsend deprivation index), compared to the least deprived fifth (OR 1.37, 95% CI 1.22 to 1.54). However, this study did not find an association between deprivation and length of PICU stay, indicating that there may be no difference in disease severity. Another study, which looked at all hospital admissions for bronchiolitis in England (not just those admitted to
intensive care), did not find an association between inpatient duration of stay and IMD at the Primary Care Trust level.\textsuperscript{15} Research from the USA, which investigated additional indicators of disease severity, including receipt of mechanical ventilation, days since breathing problems began and presence of apnoea, also found no difference between bronchiolitis severity among hospitalised infants from different median household income groups (based on ZIP-codes, which covered a median of 2,751 people at the 2010 census).\textsuperscript{85,86} Higher income groups were more likely to be admitted to an ICU, however, pointing towards a disparity in treatment based on non-clinical factors. The authors suggest that wealthier families, who can afford the “best care” may push for ICU treatment; a factor which may be less relevant in the context of the (mostly) free at the point of use health service in the UK.\textsuperscript{87}

Associations between low SEP (i.e. more deprivation), measured by parental education, income and IMD, and a doctor diagnosis of asthma in childhood, have been noted across cross-sectional and cohort studies in the UK.\textsuperscript{21,29,88} Estimates of asthma and wheeze among children in 2019 derived from the cross-sectional Scottish Health Survey are displayed in Figure 1.2.\textsuperscript{89} This chart shows a broad increase in asthma prevalence by increased deprivation, particularly where SEP is measured by the Scottish index of multiple deprivation (SIMD).

Figure 1.2. Prevalence of wheezing in the last 12 months (dashed lines) and doctor diagnosis of asthma ever (solid line) among children <16 years, by measure of SEP:

Scotland, 2019
There are also inequities in the risk of emergency hospital admissions due to the condition. A meta-analysis of 3 randomised controlled trials and 33 observational studies (mostly from North America) estimated that the odds of emergency department re-attendance was 97% greater (OR 1.97, 95% CI 1.17 to 3.34) and hospital admission 25% greater (OR 1.25, 95% CI 1.07 to 1.47) in children from low compared to high socioeconomic groups (measured by a several indicators including income, insurance status and working rank). On average, in England between 2002 and 2015, the rate of emergency hospital admissions for asthma was 3.34 times greater among people aged 5-44 years in the most deprived quintile of IMD compared to the least deprived quintile (IRR 3.34, 95% CI 3.30 to 3.38).

Inequities in the prevalence of asthma in the UK are relatively high compared to similar countries. This is shown in a paper that I wrote as part of a rotation in the first year of my PhD (the paper, published in the European Journal of Epidemiology, is included in Appendix 1.1). This study investigated the association between maternal education and reported doctor diagnosis of asthma in children ≤8 years across 10 European countries. The UK data source was the Millennium Cohort Study, which is a longitudinal cohort study of approximately 19,000 young people born across the four nations of the UK between 2000-2001. This study found that, of the 10 included countries, the UK had the second highest relative difference in offspring asthma prevalence between the highest and lowest maternal education groups. Children born to mothers with up to lower secondary level education were 73% more likely to report that their child had a doctor’s diagnosis of asthma by age 5 compared to mothers with post-secondary level education (95% CI 1.44 to 2.09), after adjustment for child sex, smoking during pregnancy, parity, maternal age and ethnicity.

Not all studies present a clear association between socioeconomic position and the prevalence of asthma. Differences in survey design, the age of the population sampled and the definition of SEP likely contribute to differing results; however, the measurement of asthma is likely the biggest contributing factor. As explained in Section 1.1.2, asthma is a descriptive label applied to diverse array of symptoms, and these also likely have a diverse array of aetiologies. It therefore follows that the influence of SEP may vary among children with different types of asthma phenotypes and, by treating asthma as one homogenous disease in research, we fail to tease out these associations. Research using the Millennium Cohort Study found a slightly stronger relationship between maternal education and early remitting childhood wheeze (relative risk ratio (RRR) of 1.48 when comparing children of mothers with no qualifications to those with a degree) than for persistent wheeze (RRR 1.31 (95% CI 1.04 to 1.65).
The association between SEP and asthma also differs in studies that consider comorbid atopy. In 2014, Uphoff et al. conducted a systematic review of studies looking at the association between SEP and asthma and allergic diseases in children and adults. Overall, they found a positive association between lower SEP (i.e. more deprivation) and asthma (OR 1.38, 95% CI 1.37 to 1.39), but the opposite association was found when the outcome was limited to atopy (OR 0.67, 95% CI 0.62 to 0.72). Uphoff and colleagues write that this finding may simply represent differences in the reporting of allergies, but the evidence for this is unclear. Another explanation is derived from Strachan’s 1989 ‘hygiene hypothesis’, which proposed that declining family size, household improvements and higher personal cleanliness were contributing to the rise in allergic diseases, especially amongst richer families. Subsequent studies have provided inconsistent evidence for this theory, with the newer “microflora hypothesis” focusing on disruption to early life microbial exposure as the pathway to allergic diseases.

1.4.2. Ethnicity and migration status (not included in Figure 1.1)

Ethnicity is another grouping that can represent a structural determinant of health. Ethnicity (or ethnic group) is defined as “the social group a person belongs to, and either identifies with or is identified with by others, as a result of a mix of cultural and other factors including language, diet, religion, ancestry, and physical features”. The term ethnic minority is used to describe people regarded by themselves or others as not being from the majority ethnic group in the population (i.e. white or white British in the UK). People from ethnic minority groups in the UK tend to experience greater disparities in socioeconomic and health outcomes (although experiences differ vastly by ethnicity). Krishna et al. suggest that poor clinical health outcomes for allergy and asthma, in particular, are due to a multitude of factors including “generally lower socio-economic status, language barriers, cultural, social, religious aspects and stigma/taboo associated with chronic disease that may impact on engagement with clinical services and compliance to therapy.”

Migration status, defined by residence in a country that is not an individual’s country of birth, often overlaps with ethnic minority status, but is not the exact same entity. There are joint and separate pathways that may operate from ethnicity, migration status and socioeconomic position to inequities in health outcomes. Prominently, the relationships between inequity, ethnic group and migration status are not heterogeneous and, as written by the Migration Observatory, “classifying people according to their ethnic origin or immigrant background is tendentious and difficult”. The complexities in unravelling the relationship between ethnicity and health outcomes is reflected in the respiratory health literature. One study of bronchiolitis admissions to hospital in England in 2008/09 observed no difference in the distribution of
ethnicity (split into white and “other” ethnicities). However, the findings results could be biased by missingness; 15% of children had no ethnicity recorded. Moreover, the broad grouping of all ethnic minority groups will hide underlying disparities relevant to some groups. The highest rates of bronchiolitis were observed among infants from black and Hispanic ethnic groups and the lowest in Asian groups in two USA studies (hospitalisations in New York and medical records in Minnesota).\textsuperscript{102,103} Between 2004 and 2007, the odds of admittance to a PICU in the UK with a primary diagnosis of bronchiolitis was lower in South Asian infants compared to non-South Asian infants (OR 0.73, 95% CI 0.63 to 0.84).\textsuperscript{84} There is no further breakdown of ethnicity beyond South Asian.

Differences in the prevalence of doctor diagnosed asthma and severity of asthma symptoms by ethnicity and immigration status have been noted in the UK. A national cohort study using Scottish administrative data showed significant differences in the rate of hospitalisations for asthma by ethnic group.\textsuperscript{104} Compared to the white Scottish population, age-adjusted rates of admission for asthma were 20-50% greater among people from a South Asian ethnicity and 30-40% lower among people with a Chinese ethnicity. These associations remained after adjustment for SIMD and country of birth. A study using Born in Bradford cohort data found a 59% increased risk of doctor diagnosed asthma among Pakistani children compared to white children at age 4 in unadjusted logistic regression analysis (OR 1.59, 95% CI 1.11 to 2.28).\textsuperscript{105} Parental immigration status was marginally associated with an increased odds of asthma diagnosis (OR 1.33 95% CI 0.97 to 1.81). However, with no pattern seen among other measures of asthma, including reported wheeze and asthma medication, the study authors concluded that “traditional risk factors such as gender, family history, socio-economic status and child’s medical history may be stronger risk factors than ethnicity or familial migration patterns.”

In a study conducted in Leicestershire, reported rates of ever having asthma, wheezing or bronchitis were twice as high among white women compared to South Asian women (21.8% vs. 10.9%).\textsuperscript{106} However, this gap closed when restricting south Asian women to those born in the UK (16.2%), highlighting the potential influence of early life exposures in the UK. This reflects findings from a linked administrative data study in Sweden, which showed a persistent decline in the odds of asthma medication with age at migration in children from a range of low or median income regions.\textsuperscript{107} The pattern was noted both among children who were adopted from abroad by a Swedish family and those who emigrated with their parents, further highlighting the role of environmental factors in asthma.
1.4.3. Geographical location

Across the globe RSV exhibits a latitudinal gradient in the timing of epidemics.\textsuperscript{108} In the Northern hemisphere this manifests as RSV activity beginning in July in tropical sites and spreading further North to reach high-latitude sites in January. A systematic analysis of global viral patterns found, in Europe, where there is more longitudinal that latitudinal variation, a later RSV onset in the East compared to the West (0.8 months across 20°E), $r = 0.45$, $p = 0.03$, but no clear association between North and South, $r = 0.11$, $p = 0.61$.\textsuperscript{108} In temperate maritime climates such as the UK, RSV epidemics occur during the colder months.\textsuperscript{109,110} Climatic factors, including temperature, humidity, precipitation and vapour pressure have all been linked to RSV outbreaks.\textsuperscript{108,110,111} A study of the infectious disease data reported to Public Health England (PHE) observed a positive correlation between air frost ($r = 0.69$), and a negative correlation between max temperature ($r = -0.60$), and number of cases of RSV infection one month later.\textsuperscript{112}

Climate change has been implicated in changes to the length of RSV epidemics among young children in the UK. A study of bronchiolitis hospital admissions and laboratory isolations of RSV in England and Wales between 1990 and 2004 found that the epidemic length was associated with a 2.5 to 3.1 week decrease per 1 degrees increase in temperature over this time.\textsuperscript{113} Alongside infectious and cardiovascular diseases, asthma and allergies are described as being “at the front line of the sequelae of climate change”.\textsuperscript{114} Cecchi et al. describe the impact of climate change on allergies in terms of the exposome—an increase in environmental exposures, such as pollen load and moulds, that cumulatively impact on an individual from birth onwards.\textsuperscript{115} Air pollution, a precursor to climate change that is socially patterned in terms of exposure,\textsuperscript{116} is another important component of this sequence (discussed further in Section 1.5.3, below).

There is some evidence suggesting that geographical location influences the development of asthma. One study of current reported asthma across the USA and Australia, found that a 10° change in latitude from South to North (approximately 220km) was associated with a 2.0% increase in the prevalence of asthma.\textsuperscript{117} In mutually adjusted analyses of the USA sample, annual mean air temperature (°C) and solar radiation (kWh/m\textsuperscript{2}/day) were negatively associated with asthma prevalence, $r = -0.42$ and $r = -0.10$, respectively. A 2016 Cochrane review of clinical trials found a “clinically and statistically significant protective effect of vitamin D against severe exacerbation of asthma”,\textsuperscript{118} p.21 further suggesting that sunlight plays a role in the disease. The relative size of the UK means there is unlikely to be substantial geographical variation in the prevalence of asthma within the country due to latitude or climate;
however, changing climatic features across the year, including temperature fluctuations and humidity have been associated with asthma symptoms and mortality across the country.\textsuperscript{119,120}

1.4.4. Built environment

I use the term built environment to refer to the complex of buildings, streets, greenspace and infrastructure that make up the physical spaces that humans live within.\textsuperscript{121} These physical spaces can contribute positively to health and wellbeing by influencing lifestyle factors such as exercise and social interactions.\textsuperscript{122,123} On the other hand, the built environment can facilitate exposure to harmful agents, such as viruses, outdoor pollution and toxins. These negative exposures are particularly important in the context of bronchiolitis and asthma. There is some evidence from non-UK countries that concentrated populations (typically found in high density built environments) affects the transmission of RSV. In Pitzer et al.’s study of the spatiotemporal dynamics of RSV in the USA, population density at the state level was positively correlated with the reproduction number of RSV ($r = 0.77$).\textsuperscript{111} In another study, at the ZIP code-level in Connecticut, population density was correlated with earlier RSV epidemic peak and higher incidence rates.\textsuperscript{124} The built environment is inherently linked to levels of outdoor air pollution with road transport sources producing two of the main pollutants associated with respiratory morbidity, nitrogen dioxide (NO\textsubscript{2}) and particulate matter less than 2.5/10 micrometres (PM\textsubscript{2.5/10}) (discussed further in Section 1.5.3).\textsuperscript{24,125} Facets of the indoor environment, particularly damp and mould, are also relevant to poor respiratory health (see Section 1.5.10).

1.4.5. Neighbourhood

SEP and neighbourhood are intrinsically linked, and in some instances are inseparable, as neighbourhood level indicators are sometimes used to define SEP, as in this study. There is evidence to suggest that measurements at the neighbourhood level are less volatile than other measures such an income.\textsuperscript{126} Of 13 studies in Uphoff et al.’s systematic review reporting on area deprivation or average household income within a country, 12 showed an association between higher deprivation and asthma or wheeze and 1 found ‘no significant result’.\textsuperscript{92} There are also risk factors for ill-health that exist at the neighbourhood level, such as physical deterioration and crime, that individual or household measures cannot characterise.\textsuperscript{127–129} Unequal access to resources may further enhance inequities (discussed further in the health system Section, 1.6).\textsuperscript{121} Moving between neighbourhoods may also have an impact on respiratory health in children. Using a cohort of 2,619 children from California, USA, Cantu et al. investigated the impact of mobility between poor and non-poor neighbourhoods.\textsuperscript{130} After a period of at least 4 years, 6.2% of the cohort moved into and 11.5% moved out of a poor
neighbourhood. Membership of the former group was associated with higher rates of childhood asthma than children who remained in a poor area at all times of the study (19.5% vs. 16.7%, 13.2% overall prevalence). The authors theorise that moving may be a sign of family instability or stress; however, the models did not account for possible mediators such as passive smoking that may help explain this association.

1.5. Intermediary determinants

For the most part, structural determinants of health do not act directly on health outcomes but, rather, work through intermediary determinants (or risk factors) to determine health. In statistical terms, these intermediary determinants are known as mediators, a concept which will be discussed further in Chapter 6. In this section, I outline how each of the intermediary determinants listed in Figure 1.1 is associated with socioeconomic deprivation and inequities in bronchiolitis/asthma. For illustrative purposes, these are grouped by timing of exposure (in-utero, perinatal and postnatal), although some may exert their influence at more than one time (e.g. smoke exposure in-utero and passive smoking in the early years). Notably, this list is not exhaustive and other risk factors have been identified, with varying levels of association noted.

1.5.1. Maternal smoking during pregnancy

Rates of smoking in the population follows a clear social gradient, and this pattern is also observed among pregnant women.\textsuperscript{131,132} The Scottish Public Health Observatory reports that in 2018/19, 26.6% of women living in the most deprived fifth of SIMD reported smoking during pregnancy compared to just 3.3% in the least deprived fifth.\textsuperscript{133} In England in 2018/19, 12.8% of women living in the most deprived clinical commission group (CCG), as measured by IMD, reported that they smoked at the time of delivery compared to 6.1% in the least deprived CCG.\textsuperscript{134,135}

Two Northern American studies have also looked at the association between maternal smoking during pregnancy and medical visits for bronchiolitis. An observational cohort study, based on term non low-birth weight infants enrolled in the Tennessee Medicaid Program from 1995 to 2003, found that maternal smoking was associated with a 14% increased risk of a bronchiolitis diagnosis (adjusted HR 1.14, 95% CI 1.10 to 1.18) and a 28% increased odds of hospitalisation for the condition (adjusted OR 1.28, 95% CI 1.20 to 1.36).\textsuperscript{136} A Canadian study, conducted on administrative records of all infants born the geographical area known as the Georgia Air Basin (British Columbia) between 1999 and 2002, found a stronger association between maternal smoking and hospitalisation for bronchiolitis (adjusted HR 1.47, 95% CI
A study of Scottish administrative data presented the association between maternal smoking in pregnancy and subsequent offspring respiratory admissions. Hazard ratios by smoking status on bronchiolitis admissions within the first year of life and asthma admissions between the 1st and 5th year are displayed in Table 1.2. Women who reported being a current smoker during pregnancy had the highest risk of subsequent offspring bronchiolitis (unadjusted HR 1.79, 95% CI 1.73 to 1.84) and asthma (unadjusted HR 1.51, 95% CI 1.43 to 1.59) when compared to women reporting never smoking.

Table 1.2. Rates of hospital admissions for bronchiolitis (birth to 1 year) and asthma (1 year to 5 years), by mothers smoking status at first antenatal booking (from Lawder at al.)

<table>
<thead>
<tr>
<th></th>
<th>Infants at risk</th>
<th>Infants with ≥1 admission</th>
<th>% admitted</th>
<th>Crude HR (95% CI)</th>
<th>Adjusted HR* (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Bronchiolitis</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><em>Never smoked</em></td>
<td>391,861</td>
<td>9,499</td>
<td>2.4</td>
<td>Reference</td>
<td>Reference</td>
</tr>
<tr>
<td><em>Current smoker</em></td>
<td>159,819</td>
<td>6,861</td>
<td>4.3</td>
<td>1.79 (1.73–1.84)</td>
<td>1.43 (1.38–1.48)</td>
</tr>
<tr>
<td><em>Former smoker</em></td>
<td>59,326</td>
<td>1,534</td>
<td>2.6</td>
<td>1.07 (1.01–1.13)</td>
<td>1.03 (0.97–1.08)</td>
</tr>
<tr>
<td><em>Unknown</em></td>
<td>85,997</td>
<td>2,505</td>
<td>2.9</td>
<td>1.21 (1.16–1.27)</td>
<td>1.15 (1.09–1.21)</td>
</tr>
<tr>
<td><strong>Asthma</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><em>Never smoked</em></td>
<td>382,327</td>
<td>3,436</td>
<td>0.9</td>
<td>Reference</td>
<td>Reference</td>
</tr>
<tr>
<td><em>Current smoker</em></td>
<td>156,123</td>
<td>2,164</td>
<td>1.4</td>
<td>1.51 (1.43–1.59)</td>
<td>1.29 (1.22–1.37)</td>
</tr>
<tr>
<td><em>Former smoker</em></td>
<td>58,025</td>
<td>608</td>
<td>1.0</td>
<td>1.16 (1.07–1.27)</td>
<td>1.10 (1.00–1.20)</td>
</tr>
<tr>
<td><em>Unknown</em></td>
<td>83,113</td>
<td>735</td>
<td>0.9</td>
<td>1.04 (0.96–1.12)</td>
<td>1.01 (0.92–1.10)</td>
</tr>
</tbody>
</table>

*Adjusted for: maternal age; infant gender, SIMD; mother’s socioeconomic status (measured by occupation); father’s socioeconomic status (measured by occupation); parity; mode of delivery, infant feeding at 6–8 weeks and country of birth.

Associational studies have also presented an association with passive smoking exposure after birth and bronchiolitis/asthma. In many cases, the size of the association is larger than that of smoke exposure in-utero. For example, a meta-analysis pooling results of 7 studies identified a 2.51 increased odds of bronchiolitis where infants were exposed to passive smoke by any household member (adjusted OR 2.51, 95% CI 1.96–3.21). However, there is no clear indication of how exposure to smoke exposure in-utero was accounted for in these studies. In another systematic review, contradictory findings were noted among studies that looked at the association between passive smoking in early years and non-atopic asthma. Worsened bronchiolitis outcomes due to passive smoke exposure have also been noted. A cohort study of children admitted to a children’s hospital in Liverpool for bronchiolitis found that infants living with a tobacco smoker had increased odds of needing oxygen supplementation and mechanical ventilation than infants without a household tobacco smoker.
1.5.2. Maternal asthma

As discussed above in Section 1.1.2 in reference to the paediatric population, the prevalence of asthma in adults also exhibits a socioeconomic gradient. In 2018, 21% of Scottish women in the most deprived quintile of SIMD reported a doctor’s diagnosis of asthma compared to 15% in the least deprived SIMD quintile. Maternal asthma, in turn, is with an increased odds of offspring bronchiolitis and asthma. A study of otherwise healthy infants who visited a health care facility for bronchiolitis found a risk 39% greater (HR 1.39, 95% CI 1.30 to 1.48) in infants of mothers with asthma compared to mothers without asthma (identified using clinical records), after adjustment for a multitude of risk factors. Analysis of data from the Oxford record linkage study showed a strong association between maternal asthma and offspring hospital admission for asthma after 2 years of age (OR 3.07, 95% CI 2.61 to 3.60 in multivariable analyses).

It has been suggested that perinatal factors, including birthweight and gestational age, may sit on the pathway between maternal asthma and child respiratory ill-health. However, the association between maternal and offspring asthma is complex and likely due to a mixture of factors including familial disposition to respiratory ill health, direct effects of maternal asthma on their offspring and shared environmental factors. In Strina et al.’s systematic review, familial history of asthma or atopy was the most consistently associated risk factor with paediatric non-atopic asthma, with the authors referencing the influence of genetics and/or shared environment. However, other studies that examine both maternal and paternal symptoms suggest that there is also an effect attributable to the mother only. For example, Schoos et al. examined the distinct association between mother and offspring asthma and atopy using 685 parent-child trios from the Copenhagen Prospective Studies on Asthma in Childhood 2010 cohort. They found a greater (but overlapping) association between mother and child asthma/persistent wheeze symptoms between 0 and 6 years, than with father and child symptoms (adjusted OR for mothers 2.11, 95% CI 1.46 to 3.05 compared to OR 1.55 for fathers, 95% CI 1.03 to 2.33). The authors write that “this suggests that maternal non-genetic factors seem to confer an added disease risk to the child, particularly in early life.”

1.5.3. Outdoor air pollution

The relationship between socioeconomic position and ambient air pollution levels is complex, varying by urban-rural status and across countries, but the evidence broadly points to inequities in exposure to air quality across England and Scotland. A 2006 report for the Department of Environment, Food and Rural Affairs summarised that there are higher relative concentrations of NO₂ and PM₁₀ in more deprived areas (measured by IMD/SIMD...
deciles) in England and Scotland. In England, sulphur dioxide (SO₂, produced during fossil fuel combustion at industrial plants) levels were also higher in areas with greater deprivation. These patterns are driven by urban areas (where more people live in deprived areas and there are far bigger populations generally) and, notably, there is wide variation within deprivation deciles.

A link between short-term spikes in air pollution and asthma symptom exacerbations has been well established in the research literature; however, ascertaining a causal role in the chronic exposure to ambient air pollution and the development of respiratory conditions has been more difficult. Research in this area is particularly limited by difficulties separating the effects of specific pollutants and in assigning the correct exposure level to individuals. Nonetheless, there is a large body of evidence showing an association between concentrations of NO₂ and PM₂₅ (measured prenatally and in early-life) and the onset of asthma in children. As highlighted in a nationwide case-control study conducted in Denmark, this is even the case where PM₂₅ levels are below the proposed European annual limit.

The evidence of a link specifically with bronchiolitis is less strong, particularly for chronic exposure to air pollution. One systematic review looked at the effect of several outdoor air pollutants on the risk of hospitalisation for bronchiolitis in children under the age of 2. The findings were inconsistent, hampered by variation between studies in measurement of study variables and the lack of cohort study designs. However, there is evidence of an association between air pollution and other early-life acute lower respiratory tract infections, particularly pneumonia. As part of the ESCAPE project, combined results of 10 European birth cohorts presented a 76% increased odds of doctor-diagnosed pneumonia by age 3 per 10-μg/m³ increase in PM₁₀ (adjusted OR 1.76, 95% CI 1.00 to 3.09) and 30% increased odds of pneumonia per 10 μg/m³ increase in NO₂ (adjusted OR 1.30, 95% CI 1.02 to 1.65). A meta-analysis of studies, the majority of which were in high-income countries, produced an estimate of 12% increased risk (95% CI 1.03 to 1.30) in acute lower respiratory tract infection occurrence per 10 μg/m³ increase in annual average PM₂₅.

1.5.4. Maternal age at delivery

Maternal age increases with higher levels of socioeconomic position. Of live births in England and Wales in 2019, 32% of mothers in the lowest category of the National Statistics Socioeconomic classification (a measure of SEP based on occupations) were less than 25 years at the time of birth compared to just 3% in the highest category. In contrast, 16% of mothers in the lowest category were 35 and older at the delivery of their child compared to 37% in the highest category. A similar pattern was observed in Scotland in 2019/20, where the most
common maternal age at first birth was 24 years among mothers from the most deprived SIMD quintile compared to 30 years in mothers from the least deprived quintile.\textsuperscript{159}

Risk of RSV-related hospital admission before the age of 3 was associated with earlier maternal age at delivery in a Scottish birth cohort.\textsuperscript{44} Children of mothers aged less than 20 at birth had a 43\% increased adjusted risk compared to mothers between 30 and 39 years old (adjusted HR 1.43, 95\% CI 1.26 to 1.32). Higher maternal age during pregnancy was associated with lower odds of offspring asthma without atopy at 7/8 years of age (OR 0.95, 95\% CI% 0.93 to 0.97) in the Avon Longitudinal Study of Parent and Children.\textsuperscript{29} Given the strong correlation between SEP and maternal age, it is thought that SEP-related risk factors, unmeasured in these studies, explain the majority of the noted associations.\textsuperscript{44,137} Although evidence for a causal association has not been shown,\textsuperscript{29} Koehoorn et al. theorise that a potential biological mechanism could be “related to maternal nutrition, health status, or stress that alters the development of the respiratory system of the fetus”.\textsuperscript{137}

1.5.5. Congenital heart disease, chronic lung disease and other early-life conditions

There are a number of early-onset chronic health conditions, which are associated with a greater risk of respiratory ill health, particularly RSV-related hospitalisation. The foremost examples of these are congenital heart disease and chronic lung disease, which are two of the conditions specifically listed (alongside other criteria) in recommendations for infants to receive prophylaxis against RSV infection.\textsuperscript{40,41} Congenital heart disease encompasses a range of defects, which develop prenatally and affect the structure of the heart.\textsuperscript{160} Congenital heart diseases are the most prevalent type of birth defects in the UK, affecting 60.9 per 10,000 births in England in 2018 according to the National Congenital Anomaly and Rare Disease Registration.\textsuperscript{161} Chronic lung disease includes some congenital conditions (i.e. those that develop prenatally), but primarily consists of bronchopulmonary dysplasia—chronic lung dysfunction related to extreme premature birth.\textsuperscript{162} Other notable early life conditions include Down’s syndrome, cystic fibrosis and neuromuscular disease.\textsuperscript{163,164}

Socioeconomic patterns of these conditions differ by the specific condition and underlying cause\textsuperscript{165,166}. Notably, conditions caused by chromosomal anomalies (including Down’s syndrome) are more prevalent in higher socioeconomic positions, which is primarily explained by the increased prevalence of these conditions at older maternal ages.\textsuperscript{166} Non-chromosomal anomalies, including congenital heart disease, tends to be concentrated in more deprived populations.\textsuperscript{165,166} It is thought that socially patterned risk factors, including maternal nutrition, alcohol, tobacco and chemical exposure likely contribute to this, although there are still many
unknowns.\textsuperscript{167} Parental decision making about terminations following prenatal identification is another noted factor in the prevalence of congenital anomalies.\textsuperscript{165,168} A study of the East Midlands and South Yorkshire congenital anomaly register found similar rates of anomalies detected prenatally across IMD deciles (after adjusting for maternal age). However, lower proportions of terminations after antenatal diagnosis meant that the most deprived areas had a 61% higher rate of live births (IRR 1.61, 1.21 to 2.15) associated with a congenital anomaly.\textsuperscript{168}

The risk of RSV-related hospital admission in children less than 3 years old was 3.4 times higher among children with a chronic condition (including chronic heart, lung and neurological conditions) born in Scotland between 2009 and 2015 (unadjusted HR 3.40, 95% CI 3.11 to 3.72).\textsuperscript{44} In England in 2008/07, infants with a congenital heart disease comprised 12.1% of those admitted to hospital with bronchiolitis in the first year of life, although only comprising <1% of the infant population.\textsuperscript{45} After adjustment for other risk factors, infants with a congenital heart disease had a 3.35 increased risk of bronchiolitis hospitalisation compared to infants without this condition (95% CI 2.92 to 3.84). Once admitted, children with congenital anomalies are also at an increased risk of complications, including secondary infections, and mortality.\textsuperscript{169}

In a longitudinal study covering the Province of Manitoba in Canada, congenital anomalies of the respiratory and circulatory system were associated with a 43% (HR 1.43, 95% CI 1.27 to 1.61) and 16% (HR 1.16, 95% CI 1.03 to 1.30) increased risk of an asthma-associated health care visit by age 6, respectively.\textsuperscript{170}

1.5.6. Birth weight and gestational age

Gestational age and birthweight are strongly related to one another, with gestational age likely on the pathway to birthweight.\textsuperscript{171} Children born into lower SEPs are more likely to be born before full term (< 37 weeks) and at a lower birthweight. Millennium Cohort Study data presents a social gradient in the proportion of children born with low birth weight.\textsuperscript{172} 10.1% of children born to mothers with no secondary school qualifications had low birthweight (< 2500g) compared to 4.5% of those born to mothers with an undergraduate degree or higher. A national cohort study of NHS records in England and Wales found that 6.7% of singleton children in the most deprived 10\textsuperscript{th} (measured by IMD) were born prematurely compared to 4.5% in the least deprived 10\textsuperscript{th}.\textsuperscript{173}

Hardelid et al.\textsuperscript{44} explored risk factors for RSV-confirmed hospital admissions in the first 3 years of life using a birth cohort created using linked Scottish administrative data. The association between RSV admission increased with earlier gestational age; at the lowest end of gestational age, children born at less than 34 weeks had an adjusted hazard ratio of 2.54 (95%
CI 2.10 to 3.06) compared to children born at term (37-40 weeks). Alongside gestational age, fetal growth is also implicated in birthweight, and babies that have fetal growth restrictions may have different health problems than those that are light due to prematurity. Small for gestational age (SGA)—commonly defined as birthweight below the 10th percentile for sex-specific gestational age—is commonly used in studies as a proxy for fetal growth. In their study of Scottish administrative data, Hardelid et al. observed an increased risk of RSV-related hospital admissions in the first 3 years of life among children born SGA after adjustment for gestational age and other factors (HR 1.14, 95% CI 1.03 to 1.25).

A meta-analysis of 13 cohort studies estimated that the risk of asthma in childhood was 16% higher in those born with a low birthweight (pooled RR 1.16, 95% CI 1.13 to 1.20). Birthweight below 2500g was associated with increased risk of asthma and wheezing disorders compared to children weighing between 2500g and 4000g at birth in the Born in Bradford study. A large cohort study of children from Denmark, Sweden and Finland estimated a 17% increased risk of hospitalisation for asthma between 3 and 18 years of age for every 1000g decrease in birthweight (RR 1.17, 95% CI 1.15 to 1.18) and 5% increased risk for each reduced week of gestational age (RR 1.05, 95% CI 1.02 to 1.09). On the other hand, two studies investigating the association of prematurity with non-atopic asthma found no relationship. There is mixed evidence on the role of SGA in asthma development. The combined Denmark, Sweden and Finland cohort study mentioned above found an association between SGA and asthma hospitalisation risk in children born at term only.

1.5.7. Delivery method

Caesarean section is a life-saving intervention when specific complications arise during pregnancy and childbirth, but has also been associated with a range of risks to the mother and child. Compared to many other intermediary factors discussed in this section, the evidence on socioeconomic differences in delivery method is less clear. Crude results suggest higher rates of caesareans in the least deprived mothers, however, this relationship changes when more nuanced analyses are used. A large study of administrative hospital data in England found no association in rates of caesarean sections by IMD after adjustment for clinical indication (including breech presentation and previous caesarean section). A study of the 2010 National Maternity Survey in England indicated a slightly raised odds of caesarean section in increasingly deprived groups (measured by IMD) when adjusted for ethnicity, maternal age and parity (OR 1.04, 95% CI 0.99 to 1.10). Analyses of the Millennium Cohort Study found an association between caesarean section and increased deprivation that differed depending on the parity status of the mother and whether SEP was defined by education or occupation. For first-time mothers, only (lower) household occupation was associated with
an increased risk, whereas for multiparous women only (lower level of) education was associated with an increased risk of a planned caesarean birth.

A cohort study from Western Australia found that, compared to spontaneous vaginal delivery, children born by elective caesarean had an 11% increased risk of admission to hospital for bronchiolitis in the first year of life, after adjustment for other risk factors, which again points to the influence of underlying clinical risk (IRR 1.11, 95% CI 1.01 to 1.23). A meta-analysis of 26 studies estimated that birth by caesarean section increased the odds of an asthma diagnosis between ages 0 and 18 by more than 20% compared to spontaneous vaginal delivery (elective caesarean, OR 1.21, 95% CI 1.17 to 1.25; emergency caesarean OR 1.23, 95% CI 1.19 to 1.26). Differences in the intestinal flora between babies born by caesarean and vaginal birth, has led to the hypothesis that immune system development lies on the pathway between delivery method and respiratory ill-health. However, two sibling control studies, a study design that enables researchers to look at the association independent of environment and genetic factors, in Sweden have found no evidence to support a causal association between caesarean section and asthma. These indicate that caesarean section is actually an indicator of other shared familial or intrauterine factors which increase the risk of asthma.

1.5.8. Number of older siblings

On average in the UK, the number of children in a family increases with socioeconomic deprivation. Berrington and colleagues investigated fertility histories in the UK using retrospectively reported information from two panel studies, the British General Household Survey and the UK Household Longitudinal Study. Among all women born between 1960 and 1969 (including those who have no children), lower levels of completed formal education were associated with a lower average completed family size. This ranged from a mean of 2.35 children among women who left school before or at 16 with no formal qualifications to 1.68 among women with a degree-level education or higher. This pattern is strongly related to the later age at entry into motherhood for women with higher levels of formal education. The presence of, particularly older, siblings increases a young child’s exposure to infection. Hardelid et al. estimated that, in a Scottish setting, RSV hospital admissions in children under 3 years old could be reduced by up to 34% by removing the risk posed by older siblings (population attributable fraction 34.0%, 95% CI 31.0 to 36.9). A study of RSV hospitalisations among children under 2 years in Denmark found that this association persisted after adjustment for a number of other risk factors including childcare.
The relationship between the number of siblings and asthma is less clear. As with many studies of asthma, the heterogeneous definition of asthma used by different studies likely affects the comparability of results. More than one older sibling was associated with early transient wheeze in a birth cohort study from the Netherlands (OR 1.31, 95% CI 1.07 to 1.60). However the outcome of transient wheeze, which peaked at ages 1 and 2, may be indicative of respiratory infection rather than asthma. In Strina et al.’s systematic review of the literature, six of seven studies presented no association between family size or birth order and non-atopic asthma/wheeze. The final study, which reportedly found a protective effect of siblings, was from rural Ecuador, and the results may not be applicable to a UK-setting. Using International Study of Asthma and Allergies in Childhood (ISAAC) data on 13-14 year olds from 16 affluent counties, different trends were observed based on the outcome studied. Each additional older sibling was associated with an 8% greater odds of reporting severe wheeze in the last 12 months (OR 1.08, 95% CI 1.04 to 1.12), whereas the opposite effect was shown for asthma ever (OR per additional sibling 0.96, 0.94 to 0.99).

1.5.9. Breastfeeding

To provide the ideal nutrition for the healthy growth and development of infants, the WHO recommends that babies receive breast milk exclusively for the first 6 months of life. The latest national estimate of rates of breastfeeding, from 2010, placed the UK as having one of the lowest breastfeeding rates in the world. The (now discontinued) Infant Feeding Survey reported that 34% of women were still breastfeeding their child at 6 months in 2010. This figure followed a social gradient, from 31% of women in the most deprived IMD quintile to 40% in the least deprived IMD quintile. There is evidence suggesting that this pattern very much persists a decade later. Public Health Scotland (PHS) reports that, in 2019/20, 43.9% of women were breastfeeding at their 6 to 8 week child health review, an estimate that ranged from 29.6% for women living in the most deprived Scottish IMD quintile to 63.8% in the least deprived quintile.

Breastfeeding has broadly shown to be associated with lower risk of early respiratory infections and, to a lesser extent, asthma, although there are differences based on the definition of breastfeeding used. There was an increased risk of hospitalisation for bronchiolitis among infants in the Georgia air basin region of Canada, where mothers had not initiated breastfeeding in the hospital (unadjusted HR 1.86, 95% CI 1.61 to 2.16). Pooled results from 3 hospital-based studies rated high quality in a systematic review showed a 2.24 increased odds of hospitalisation for pneumonia and bronchiolitis in under 5s that had not been breastfed at all (OR 2.24, 95% CI 1.56 to 3.20). In a Dutch population-based cohort study, breastfeeding at 6 months or longer was associated with a reduced risk of offspring
bronchitis or pneumonia up to age 4 years, compared to no breastfeeding (OR 0.63, 95% CI 0.46 to 0.87). Similar but slightly lower effect estimates were produced for breastfeeding less than 3 months (OR 0.75, 95% CI 0.56 to 1.02) and 3-6 months (OR 0.72, 95% CI 0.48 to 1.06), compared to no breastfeeding.

In a meta-analysis of wheeze trajectory-specific risk factors in children, any breastfeeding was associated with a lower odds of early transient wheeze symptoms, compared to never/infrequent wheeze (pooled OR 0.86, 95% CI 0.77 to 0.95). This association was not apparent in children with other trajectories of wheeze. Seven studies included in Strina et al.’s systematic review looked at the relationship between breastfeeding and non-atopic asthma. Of these, five found a reduced risk from breastfeeding (three looked at no breastfeeding compared to any amount of breastfeeding, one at <3 vs. ≥3 months and one at <20 vs. ≥20 weeks) and two did not report an association.

1.5.10. Indoor living conditions

Components of the indoor environment that may increase a child’s susceptibility to respiratory illness include dampness and mould, residential overcrowding and indoor air pollution (produced by cigarette smoke, heating or cooking). These housing conditions are more common among poorer households in the UK. The English Housing Survey, a national cross-sectional survey commissioned by the Ministry of Housing, Communities and Local Government (MHCLG), collects data about housing conditions at private addresses in England. This survey estimated that, in 2018, 7.2% of households in the lowest quintile of income did not have a reasonable degree of thermal comfort (related to insulation and heating efficiency) compared to 3.7% of households in the highest quintile. In addition, 5.1% of the lowest income quintile households were categorised as having “any damp” compared to 1.3% of households in the highest income quintile. It has been proposed that damp indoor air conditions prolong virus survival, which may lead to a higher likelihood of RSV infection. Persistent indoor dampness provides conditions in which microbial pollution, particularly mould, can grow. The government’s bedroom standard indicator (used in the English Housing Survey) defines household overcrowding by those with fewer bedrooms than the number needed based on age, sex and marital status of the occupants. According to the latest survey results (combined over the last 3 years to enable reliable estimates), 5.1% of households in the lowest socioeconomic group were overcrowded compared to 1.6% in the highest group (defined by occupations). Moreover, overcrowding in England was at the highest recorded figure since records began (in 1995/96) for both the social rented (8.7%) and the private rented sector (6.7%).
In their 2009 report on dampness and mould, the WHO wrote that “there is sufficient evidence of an association between indoor dampness-related factors and a wide range of respiratory health effects...including asthma development, asthma exacerbation, current asthma, respiratory infections”\textsuperscript{202 p.27}. A meta-analysis published in 2010 found “moderate but statistically significant” associations between residential dampness and mould and respiratory tract infections and bronchitis\textsuperscript{199}. In pooled analyses, Fisk et al. estimated that children living in residencies with dampness and mould had a 48% increased odds of respiratory tract infection (unadjusted OR 1.48, 95% CI 1.34 to 1.62). The proportion of the risk in respiratory infection attributable to dampness and mould in this group was estimated to be between 8.8-19.4% after adjustment for age, sex, smoking exposure and SEP. A singular study from Wellington, New Zealand using an unmatched case–control design estimates that if all housing were free from damp and mould, then 19% of these types of hospital admission for any acute respiratory infection in under 2 year olds would be prevented\textsuperscript{206}. In a systematic review published in 2012, visible mould was associated with a 49% increased odds of asthma (OR 1.49, 95% CI 1.28 to 1.72) and 68% increased odds of wheeze (OR 1.68, 95% CI 1.48 to 1.90) in children aged between 0 and 15\textsuperscript{207}.

A systematic review of 20 studies on residential crowding and laboratory-confirmed RSV hospitalisations among children younger than 5 years found a consistent increased odds of disease across different settings\textsuperscript{208}. It is thought that crowding can facilitate the spread of RSV through close contact conditions. However, the authors also offer caution when interpreting results from these observational studies, pointing out the difficulty in separating the effect of residential crowding from other socioeconomic-related risk factors such as indoor pollution. Another study used the Growing Up in New Zealand longitudinal child cohort study to look at a range of home environment risk factors in the risk of hospitalisation for acute respiratory infection during the first five years of life\textsuperscript{198}. In unadjusted analysis, associations were found between household crowding (2+ versus <2 on the crowding index, HR 1.63, 95% CI 1.42 to 1.88), as well as no heating in the house, rented tenancy (as opposed to owner), cigarette smoking in the house, dampness of the house, and heavy condensation mould or mildew in the room where the child sleeps at night. In models adjusted for a range of child, maternal and environmental factors, there were notable associations between hospitalisation for acute respiratory infection and the use of a flued or unflued gas heater in the child’s bedroom (flued HR 1.69, 95% CI 1.21 to 2.36; unflued 1.68, 95% CI 1.12 to 2.53) or gas heater as the sole type of household heating (HR 1.64, 95% CI 1.29 to 2.09). See section 1.5.1 for information about passive smoking.

1.6. The Health system
In my framework, the health system is placed just outside the other intermediary determinants of health to highlight the unique role it plays in determining the equitability of health outcomes. Access within the health system not only directly impacts on health outcomes, but it can also address differences in the exposure of other intermediary determinants and vulnerability to their effects. Understanding social determinants of health within the context of the UK health system is particularly important in my thesis, as it is through this system that the two health outcomes of interest are measured.

The national health system in the UK, the NHS, is designed so that there is equal and free access for all UK residents. In fact, economic analysis of English hospital care has shown that spending is highest amongst people living in the most deprived areas of the country. However, rather than evidencing equity, increased hospital visits may be indicative of health seeking behaviour of those in a lower SEP. In general, poorer patients tend to present symptoms later to health care professionals, are less likely to avail of preventative NHS care and report worse patient experience than their richer counterparts. The inverse care law, coined by Tudor Hart in 1971 to refer to the phenomena of good medical care varying inversely with the needs of the population it was serving, is, as put by Michael Marmot, “as relevant as ever.”

There is evidence suggesting that differences in access and uptake of preventative health care access can affect the respiratory health of very young children in the UK. Hardelid et al. found an association between delayed vaccinations at 6 months (a proxy measure for poor access to preventative health services) and risk of a RSV-related hospital admission before the age of 3 in Scotland (adjusted HR 1.14, 95% CI 1.03 to 1.25). The impact of preventative health care is particularly apparent in asthma outcomes. Asthma is considered an ambulatory care sensitive condition, which is a term used to describe conditions with symptoms that ought to be preventable by interventions in primary care. Using Hospital Episode Statistics from 2002 to 2015, Gupta et al. found a social gradient to hospital admissions with a primary diagnosis of asthma in England, indicating a similar inverse gradient to preventative care uptake. People aged 5-44 years in the most deprived quintile of IMD were over three times more likely to have an emergency hospital admission for asthma compared with the least deprived quintile (IRR 3.34, 95% CI 3.30 to 3.38).

There is also evidence of disparities in service provision for bronchiolitis and asthma once healthcare is sought. These differences are perhaps most clearly illustrated by variation in healthcare use and treatment by geographical location. Two studies have found significant variation in admission rates for bronchiolitis across hospitals in England, and concluded that healthcare provider factors, particularly admission thresholds, are likely the biggest
contributors to the variation in admission rates noted.\textsuperscript{4,215} A survey of 1,009 GPs in England in 2016 found that 19% prescribed oral corticosteroids for bronchiolitis despite NICE guidelines stating that this is not a recommended treatment.\textsuperscript{215} Other studies have shown that the availability of primary care, particular out-of-hours GP services, in England is associated with visits to the emergency department.\textsuperscript{216,217} Using cross-sectional data from English practices in 2010-2011, Fleetwood et al. showed that a 10% increase in access (measured by a composite score) was associated with a 32.1% reduction in emergency admissions for asthma across all patient ages.\textsuperscript{217} In another study, between April 2011 and March 2012, GP surgeries with the lowest levels of access (ranked using patients’ reported ability to schedule appointments from national GP Patient Survey data) had a risk of an unplanned hospitalisation for asthma in patients under 15 years more than twice than those with the highest levels of access (RR 2.22, 95% CI 2.08 to 2.38).\textsuperscript{216}

1.7. Biological factors

There are several individual-level risk factors that increase the risk of both bronchiolitis and asthma but are not necessarily affected by SEP or other structural determinants of health. These non-socially determined risk factors are presented in the conceptual model under the broad (and loosely fitting) heading of biological factors (Figure 1.1).

1.7.1. Age

Young age is the factor most strongly predictive of the risk of hospital admission for bronchiolitis. Bronchiolitis hospital admissions peak between 1 and 4 months of age and two thirds of all admissions for bronchiolitis are in children less than 6 months old.\textsuperscript{6,45,218–220} In comparison, there are very few bronchiolitis admissions after the first year of life. Between mid-2007 and mid-2012 in England, only 7.4% admissions with a primary diagnosis of bronchiolitis were amongst children 1 year and older.\textsuperscript{6} There is also a strong age-related pattern to the risk to asthma. Although, as mentioned in Section 1.1.2, it is advised that asthma is not diagnosed in young children, clinical symptoms of wheeze are frequently seen in children in the first few years of life.\textsuperscript{20,24} Additionally, differences in the timing of symptomology before the age of five have been shown to be associated with different risk factors for asthma.\textsuperscript{24} The progression of asthma is also not uniform across children. The European White Lung book estimates that of all children with asthma: about one third have persistent asthma through to adulthood; about one fifth have remission in their early teens, but see a re-appearance of asthma in adulthood; and in the remaining children (over two fifths of those with asthma), the disease remits in early teens and does not reappear.\textsuperscript{20} Although less common, asthma can
also emerge for the first time in adulthood. Occupations that expose workers to airborne agents such as chemicals and detergents have been associated with adult-onset asthma.\textsuperscript{20}

1.7.2. Season of birth

In England, infants born in September or October make up a higher proportion of admissions for bronchiolitis compared to children born in other months of the year.\textsuperscript{45} These birth months are approximately 2 months before peak viral activity, and it is suggested that a natural trough in antibody availability (alongside general immaturity of the lungs) in the first 2-4 months of life may leave these infants more susceptible to the chronic effects on the airways.\textsuperscript{53} In addition, as the NHS recommends seeking help sooner for infants under 3 months who have a temperature, this likely leads to more admission for this age group in the first place.\textsuperscript{221} An association between timing of birth in relation to winter virus peak and asthma development has also been noted.\textsuperscript{53} Wu et al. found, in a large cohort from Tennessee, USA, that children born ~4 months before the winter RSV peak had a 29% increased risk of developing asthma at 4 to 5.5 years (defined using prescriptions and hospital diagnoses data) compared to those who were 1 at viral peak (adjusted OR 1.29, 95% CI 1.19 to 1.40).

1.7.3. Sex

Clear sex differences exist in the prevalence of respiratory illness in children. There are prenatal sex differences in lung maturation growth, with females showing earlier foetal breathing, increased surfactant production and lower air resistance than males before birth.\textsuperscript{154,222} Accordingly, males have higher rates of bronchiolitis, and asthma/wheeze in the first 10 to 15 years of life, following which females begin to have an increased risk of asthma compared to males, possibly due to pubertal changes.\textsuperscript{25,93} The risk of hospitalisation for bronchiolitis amongst infants was approximately 50% higher for boys compared to girls in England and Scotland between 2000 and 2012.\textsuperscript{52} At age 10.7, 13.9% of boys were reported to have a doctor diagnosis of asthma in the last year compared to 10.1% of girls in the Avon Longitudinal Study of Parent and Children.\textsuperscript{25} Males continued to have excess asthma until 16.5 year of age when the prevalence in girls exceeded that of boys (12.6% versus 11.9%).

1.7.4. Genetic risk of bronchiolitis and asthma

Genetic background has been proposed as a contributing factor to explaining why only a proportion of infants, most with no underlying conditions, develop bronchiolitis when the majority of the population are infected with RSV by age 2.\textsuperscript{223,224} However, research findings on genetic risk factors of bronchiolitis is limited and no evidence has so far been produced from Genome-wide Association Studies (GWAS)—described as the preferred method for
gene discovery. Based on the results of twin heritability studies, it is through that asthma is a more heritable condition than bronchiolitis, with an estimated genetic contribution of about 50% compared to less than 20%. In addition, GWAS have been used to identify several genes that are associated with asthma. However, only a small proportion of asthma heritability has so far been explained by these genetic variants.

Moreover, genetic factors cannot explain the sudden rise in the prevalence of asthma at the end of the 20th Century. Interactions between genetics and environmental risk factors, alongside epigenetic exposures, have been proposed to explain differences in disease susceptibility and severity. Epigenetics describes modification of gene expression that can then be passed down to offspring, and offers advancement in understanding asthma susceptibility, although, as warned by Meyers et al. "caution must be exercised where the direction of causality is unknown, as epigenetic marks could be either a cause or consequence of disease". As with non-genetic studies of asthma, research has been hindered by the use of one simple measure of asthma, rather than considering the range of phenotypes under this umbrella condition.

A prominent theory in the bronchiolitis-asthma causality debate is that there is a shared genetic predisposition to both conditions. A Danish population-based twin study demonstrated that genetic determinants of severe RSV infection overlap with asthma. Other studies have looked at asthma and atopy within families, and found that family history of asthma or atopic disease is associated with childhood acute lower respiratory tract infections, further indicating a genetic component across these diseases. Candidate gene association studies have previously identified some immune response genes implicated in both bronchiolitis and asthma, although only a handful of common polymorphisms have been identified so far from GWASs.

1.8. Public health policy and practice

In this section, I will introduce current and planned English and Scottish public health policy and practice related to the prevention of bronchiolitis and asthma, with particular reference to attempts to lessen inequity in the rates of these two conditions. Notably, although England and Scotland both reside under the same sovereign state (the UK), health and social services fall under one of many devolved matters. Since the Scotland Act 1998, the NHS, funding, health regulation and public health, amongst other aspects of health care in Scotland, are the responsibility of the Scottish Parliament. PHE has been responsible for improving the population’s health and addressing inequalities in England since its inception in 2013.
Scottish equivalent, PHS, was formed in April 2020, taking over the function of several pre-existing public health bodies.\textsuperscript{234}

I write about the implications of the SARS-CoV-2 pandemic in my discussion chapter.

1.8.1. RSV infection and bronchiolitis

Infection control

Public guidance on RSV infection by PHE was published in 2008.\textsuperscript{235} This document gives brief information about transmission, at risk groups, symptoms and diagnosis. It has one preventative statement, ‘transmission can be prevented through standard infection control practices such as hand washing’. A 2011 Cochrane review concluded that simple and low cost interventions such as handwashing were effective at reducing the spread of respiratory viruses, especially around younger children.\textsuperscript{236} However, there is limited research on the population impact of such campaigns.\textsuperscript{237} An evaluation of the national CleanYourHands campaign—a Department of Health funded campaign to improve hand hygiene in hospitals in England and Wales—showed a reduction in some infection rates, although a causal association cannot be assumed from the study design.\textsuperscript{238} Health Protection Scotland, the public health domain previously concerned with health protection in Scotland (before PHS was created), reports on seasonal respiratory pathogens without specifically referencing prevention.\textsuperscript{239}

Palivizumab prophylaxis

PHE guidance on RSV infection mentions the lack of treatment available for RSV infection, although palivizumab is cited as a preventative therapy that is available for infants at high risk of infection.\textsuperscript{235} The risk factors that infer eligibility for palivizumab (chronic lung conditions or congenital heart disease, with preterm birth) are more common in low SEP groups,\textsuperscript{168,172} therefore we may expect this preventative treatment to reduce inequity in hospital admission rates to some degree. However, as only about 15% of infants admitted for bronchiolitis have one or more known predisposing risk factors, and a minority of these are eligible for palivizumab, this is unlikely to have a big impact on overall inequity.\textsuperscript{45} Notably, there is no national record of access to palivizumab in England, making it difficult to assess the potential impact of this drug. The limited research points to a large variation between hospital trusts, and limited uptake of this preventative medication overall. Using a linked administrative dataset, Zylbersztejn and Hardelid estimated that less than 20% of eligible infants were
actually prescribed palivizumab across 43 acute hospital trusts in England between 2012 and 2016.\textsuperscript{240}

\textit{Vaccination}

RSV vaccines are cited as the most promising avenue for the prevention of bronchiolitis,\textsuperscript{15} with research and development for this vaccine is listed as a strategic priority for the WHO.\textsuperscript{241} It is hoped that maternal immunisation against RSV will protect offspring during the first few months of life, although results from Novovax’s phase 3 randomised controlled trial (PREPARE) in early 2019 did not meet this primary endpoint.\textsuperscript{242,243} Promisingly, GlaxoSmithKline begun a phase 3 study of another RSV maternal candidate vaccine in November 2020.\textsuperscript{42} Once a RSV vaccine is available, there is still the barrier of immunisation uptake, which has typically been lower among pregnant women compared to other populations. For example, since a national outbreak of pertussis (whooping cough) that led to 14 deaths in babies under 3 month olds in 2012, immunisation for pertussis has been offered to all pregnant women by the NHS.\textsuperscript{244} However, according to statistics published by Public Health England, around 70% of women took up this vaccine in 2018/19. The influenza vaccine, which has been recommended in this population since 2010, had an uptake of only 45% among pregnant women in England during the 2018-19 flu season.\textsuperscript{245} This is compared to an average coverage of 90.9% for the 9 routine childhood vaccinations in England over the same period.\textsuperscript{246}

Childhood immunisation statistics point to further differences by level of deprivation. Figures published by the Scottish Public Health Observatory, for example, show that there is a high uptake for all vaccines routinely recommended for children by 24 months of age, but that rates follow a social gradient.\textsuperscript{247} At year-end 2019, children aged 24 months in the least deprived SIMD quintile had an average uptake of 95.8% across the four recommended vaccines compared to an average of 91.9% in the most deprived quintile. This suggests that whilst a RSV vaccination will significantly reduce overall bronchiolitis rates, it is unlikely to decrease inequity (and may even increase it) if current methods to encourage uptake are used.

Awareness of RSV and bronchiolitis is important when implementing preventive interventions, particularly given the reported decline in childhood vaccination uptake in recent years.\textsuperscript{246} A study of pregnant women at 4 hospital sites in Southern England found that, whilst only 14% reported never hearing of bronchiolitis, 71% reported never hearing of RSV.\textsuperscript{248} 36% of midwives, also surveyed in this study, had also never heard of RSV. Women who were more familiar with RSV said that they were slightly more likely to accept RSV vaccination if routinely recommended compared with women with no understanding of RSV. ‘More than a cold’ is a
campaign run by the biopharmaceutical company, AbbVie, in conjunction with several children’s charities to raise awareness of bronchiolitis, especially amongst parents.\textsuperscript{249} The impact of this campaign is unclear.

1.8.2. Asthma

NHS England states that childhood asthma is the most common chronic condition amongst young people in the UK, and the number one reason for paediatric emergency hospital admissions in England.\textsuperscript{250} In 2011, the Department of Health set out an outcomes strategy for chronic obstructive pulmonary disease (COPD) and asthma in England.\textsuperscript{251} This strategy states that “the causes of asthma are not well understood, so prevention of asthma is not currently possible” and the focus of recommendations are on accurate diagnosis, treatment and symptom control. The National review of asthma deaths, published by the Royal College of Physicians in 2014, similarly focused on poor control of asthma symptoms, inappropriate prescribing and patient monitoring.\textsuperscript{33} The English Quality and Outcomes Framework, a voluntary incentive program for GP practices, includes several indicators intended to improve the care of patients with asthma.\textsuperscript{252} Specific quality indicators include maintaining an asthma register, updating patients’ symptoms and co-morbidities and offering support/treatment to current smokers. Whilst socioeconomic deprivation is associated with severe asthma symptoms and, therefore, clinical factors are likely an important focus in reducing inequity in outcomes of people with asthma, the focus of this PhD is on preventing asthma in the first place.\textsuperscript{69}

\textit{All our health} is a framework of evidence that was published in 2015 by PHE, “to guide healthcare professionals in preventing illness, protecting health and promoting wellbeing.”\textsuperscript{253} One section of this report is related specifically to respiratory disease. This document refers to inequality in avoidable mortality due to respiratory disease and the cost these conditions have on the health system and wider economy. In 2015, the website \textit{Respiratory Futures} was set up by NHS England in collaboration with the British Thoracic Society to provide professionals with local area-level resources related to respiratory health.\textsuperscript{254} Smoking is cited as a preventative priority for almost all sustainability and transformation partnerships in England; a factor also frequently mentioned in the Department of Health’s 2011 COPD and asthma outcomes strategy.\textsuperscript{251} Other priorities highlighted in \textit{Respiratory Futures} include integrated services, early diagnosis and reducing emergency admissions.\textsuperscript{254} Leicester, Leicestershire and Rutland Sustainability and Transformation Partnership are specifically acknowledged for their work in tackling the wider determinants of the health through improved transport, air quality and housing.
Good Places Better Health for Scotland’s Children was part of the Scottish Government’s 2008 strategy on health and environment. This strategy, which focused on the role of creating health environments, considered asthma as one of four major health challenges facing children in Scotland. As part of this strategy a health evidence assessment was conducted on asthma in 2012, which reviewed the risk factors for asthma and potential actions to reduce the scale of the problem. Socioeconomic factors are cited in this assessment as an influence in the prevalence of asthma, with inadequate housing and smoking mentioned specifically in relation to deprivation. Two related actions emerged from expert workshops: “educational initiatives on smoking should acknowledge socio-economic, stress and social class contexts”; and “address socio-economic related smoking pressures”. The outcome of this work is unclear, and no clear continuation of the strategy documented.

1.8.3. Looking forward

The NHS (England) Long Term Plan was first published on 7 January 2019. This report outlines the plan for the NHS in England over the next decade. It references respiratory conditions as an area where the NHS will be intensifying its focus, citing that lung conditions cost wider society £9.9 billion annually. Section 3.83 of the NHS Long Term Plan refers specifically to social deprivation and respiratory disease:

“Incidence and mortality rates for those with respiratory disease are higher in disadvantaged groups and areas of social deprivation, where there is often higher smoking incidence, exposure to higher levels of air pollution, poor housing conditions and exposure to occupational hazards.”

The NHS Long Term Plan cites specific areas of change needed to reduce incidence of and improve outcomes for respiratory disease, including reductions in hospital stays due to these conditions, tackling risk factors that contribute to higher rates in disadvantaged groups, improved detection and diagnosis, and pulmonary rehabilitation. In addition, the ‘best start in life’ is one of the ten priorities listed in the PHE strategy 2020 to 2025. This report also notes that social disadvantage in health is persistent and in some cases, the gap is widening. Two particularly pertinent actions set out to prevent health inequalities are specialist smoking cessation services for pregnant women and commitment to lower air pollution.

NHS Health Scotland published a five-year Strategic Framework in 2017 entitled A Fairer Healthier Scotland 2017-22. This framework has a strong focus on prevention and reducing health inequalities across the board. Relevant particularly to bronchiolitis and asthma is the commitment to reduce rates of smoking, increase the proportion of infants with a healthy
birthweight and improve suitable housing options. Putting this strategy into action, the *NHS Health Scotland delivery plan for 2019-20* includes specific short term goals to improve Scottish health.\(^{260}\) These commitments include: progressing the Scottish Burden of Disease Study to identify risk factors for health inequality; supporting the implementation of the tobacco control action plan; and providing expert input into the development of strategies that aim to improve health and reduce inequalities for children, young people and families.

### 1.9. Chapter summary

To conceptualise the potential pathways to inequity in bronchiolitis and asthma rates in the UK, this chapter presents findings from the literature on the association between SEP, bronchiolitis and asthma using a social determinants of health framework. Through this framework, individual socioeconomic position is viewed as a structural determinant of health that, along with other structural factors including geographical location and the built environment, influences exposure to intermediary determinants of health. The intermediary risk factors discussed are focussed on the pre- and post-natal period and include tobacco smoke and air pollution exposure, premature birth, chronic conditions and housing conditions. Together with non-modifiable factors, such as age, sex and genetic risk, intermediary determinants ultimately modify the likelihood that a child will develop bronchiolitis and/or asthma. Current policy around bronchiolitis and asthma in England and Scotland is focussed, for the most part, on treating the symptoms of the conditions although future strategies place emphasis on preventative risk factors (particularly of reducing tobacco smoke exposure) and reducing socioeconomic inequity.
Chapter 2. Rationale, aims and thesis structure

Chapter overview

In this chapter, I summarise the key findings and future priorities that have emerged from the literature presented in Chapter 1. These form the justification for my work. I present the rationale for my PhD project, followed by my specific aims and objectives. I will finish by outlining the structure of this thesis.

2.1. Literature summary

2.1.1. Key findings from Chapter 1

- Bronchiolitis is the most common cause of hospital admission among infants and asthma is estimated to affect approximately 10% of children by their mid-teens in England and Scotland.\(^{18,261}\)
- In line with the seasonal patterns of RSV, which is the dominant pathogen that causes bronchiolitis infections, the burden of bronchiolitis on NHS hospitals is substantial in winter months.\(^{4,6}\)
- Asthma is estimated to cost the NHS around £1.1 billion per annum, including 85,000 hospital admissions and 108,000 GP consultations each year.\(^ {32}\)
- It has been established that hospitalisation with RSV-related bronchiolitis in infancy is associated with an increased risk of developing wheeze and asthma in childhood, but there is ongoing debate about the nature of this association.\(^ {49–53}\)
- There is a socioeconomic gradient in the incidence rate of hospitalisation for bronchiolitis and the prevalence of asthma in England and Scotland.\(^ {4,82,93}\)
- Alongside socioeconomic factors, wider structural factors implicated in the prevalence of these respiratory conditions include geographical location and the associated climate, as well as the built environment, particularly in relation to air pollution.\(^ {24,108,262}\)
- A multitude of socially patterned prenatal and perinatal risk factors have been shown to have an association with the development of bronchiolitis and asthma, including tobacco smoke exposure, congenital anomalies and prematurity.\(^ {44,138,263}\)
- Other non-socially patterned risk factors for bronchiolitis and asthma include child’s age, sex and, to a lesser extent, genetic variation.\(^ {5,24,25,82,225,226}\)
• The health system in the UK is designated free and equal for residents and there are national clinical guidelines in place for the treatment of bronchiolitis and asthma; however, there are disparities in the uptake of care, particularly preventative treatments, and in service provision across the UK.

• Preventative treatment of bronchiolitis is primarily focussed on the development of a RSV vaccine, but this programme is some years away from implementation and it is not currently clear whether this will tackle inequity in bronchiolitis incidence.

• In the past, asthma policy has been concentrated on treatment through improvements in symptom management and more streamlined and effective clinical pathways; however, more recent policy documents appear to be introducing a new focus on reducing inequalities and wider preventative action.

• The NHS Long Term Plan for England cites specific areas of change needed for managing and preventing respiratory disease, including reductions in hospital stays due to respiratory conditions and tackling risk factors that contribute to higher rates in disadvantaged groups.

• NHS Scotland is similarly concerned with inequalities in health outcomes, particularly childhood asthma, and is taking active steps to understand risk factors for disease burden.

2.1.2. Priorities driving my work

• Without a good understanding of the longitudinal pathways to bronchiolitis and asthma, it is difficult to design effective policies for prevention.

• There is a wealth of research identifying factors that are associated with an increased risk of bronchiolitis and asthma; however, a major limitation of the presented evidence lies with how risk factors and their interrelated pathways are considered in adjusted models.

• Methodological constraints have further hampered efforts to get unbiased causal estimates of disease risk.

• It is well established that early-RSV infection (and therefore bronchiolitis) is associated with an increased risk of asthma but the reasons for this remain unclear.

• There has been a lack of joined up work to understand the common or related pathways in the development of both of these conditions.

• Furthermore, research on the link between RSV and asthma has not considered the common role of socioeconomic deprivation as a risk factor for both conditions.

• Investigating inequities in hospital admissions for bronchiolitis, in particular, will offer guidance on the planning of future RSV vaccines.
• Through the use of more sophisticated methodology that can disentangle these complex relationships, we can work out how and when to intervene to prevent either or both of the conditions.

2.2. Thesis rationale

Bronchiolitis and asthma are major causes of morbidity among children in the UK and are responsible for substantial burden on the national health care system.\textsuperscript{4,32} Despite this, there are currently no preventative or curative measures for the majority of children that develop bronchiolitis or asthma.\textsuperscript{15,47} There is a large body of evidence investigating environmental, social and biological risk factors for these conditions, but much of this work does not fully consider the ways in which various risk factors interact with each other (and potentially bias results). There is also a myriad of research looking specifically at the link between RSV bronchiolitis and subsequent asthma,\textsuperscript{53,56,264} but precise risk factors leading to both conditions remain elusive. What we do know is that a prevailing facet of both conditions is the inequity in prevalence across socioeconomic groups.\textsuperscript{4,29} Identifying the factors that may mediate or confound the pathway between socioeconomic deprivation, bronchiolitis and asthma will help to unravel this complex picture. This is a pertinent step to working out how and when to best intervene to prevent either or both of the conditions. By focussing on early life risk factors, the aim is to reduce the burden of bronchiolitis and asthma by closing the gap in the incidence/prevalence of these conditions between children born into the most and least deprived socioeconomic groups.

2.3. Thesis aims and objectives

The aim of this project is to describe associations and explore pathways between early life socioeconomic deprivation, infant bronchiolitis and asthma in mid-childhood using administrative health datasets. Overall, this PhD will inform where interventions are best placed to reduce the burden of childhood respiratory ill health by closing the gap between the most and least deprived socioeconomic groups.
Chapter 2 - Rationale, aims and thesis structure

| **Objective 1** | To describe the association between seasonality, early life socioeconomic position and bronchiolitis admission rates among infants across England. |
| **Objective 2** | To investigate the mediating contribution of maternal smoking in the pathway between early life socioeconomic position and bronchiolitis admission rates in infancy. |
| **Objective 3** | To describe typical trajectories of asthma/wheeze among children <10 years in Scotland and their association with early life risk factors, including socioeconomic position. |
| **Objective 4** | To investigate the mediating contribution of bronchiolitis in the pathway between early life socioeconomic position and mid-childhood asthma/wheeze trajectory groups. |

2.4. Thesis structure

In Chapter 1, I presented a conceptual framework based on the Commission on Social Determinants of Health (CSDH) that differentiates between the structural and intermediary determinants of ill health. Within this framework, I have presented the current evidence on risk factors relating specifically to bronchiolitis and asthma in a UK context. I have highlighted areas of policy and practice in England and Scotland that are relevant to preventing and reducing inequity in bronchiolitis and asthma rates.

In Chapter 2, I have outlined my rationale as well as the overarching aims and objectives of my PhD.

Chapter 3 introduces the administrative datasets and derived birth cohorts that I use in this thesis. I will outline the various roles I have undertaken as part of my PhD, including applying for research approval and cleaning datasets. Lastly, I will discuss the implications of patients opting out of sharing their data with application to maternity information and research.

In Chapter 4, I present the first of two studies that focus on the seasonality of bronchiolitis admissions in England. Using a birth cohort created from administrative hospital data, I explore spatial variation in the seasonality of hospital admissions for bronchiolitis and its association with local demographic characteristics.

Chapter 5 uses the same English longitudinal dataset as in Chapter 4 to quantify the extent to which seasonality of bronchiolitis admissions is modified by socioeconomic position at the
individual level. For a wider understanding of the burden of bronchiolitis on the English NHS, I also calculate the proportion of all infant hospital admissions in England attributable to bronchiolitis by year, month and socioeconomic group.

Using a causal inference framework, Chapter 6 tackles objective 2. I use a birth cohort constructed from Scottish administrative data to quantify the mediating effect of maternal smoking during pregnancy on the pathway between socioeconomic deprivation and hospital admission for bronchiolitis. To guide the selection of confounders in my model, I apply an evidence synthesis for constructing directed acyclic graphs.

In Chapter 7, I model trajectories of asthma/wheeze symptoms among children aged between 2 and 10 in Scotland and describe their association with early-life risk factors. I then use these trajectory groups as the outcome in mediation analysis to explore the mediating contribution of bronchiolitis in the pathway between early life SEP and mid-childhood asthma/wheeze.

Chapter 8 pulls all the work in this thesis together into a summary chapter. I highlight the main findings from my research and discuss the implications of the work, including relevance to public health policy and practice.
Chapter 3. Introduction to thesis datasets

Recap and chapter overview

Chapters 1 and 2 covered an overview of previous literature in the area, the conceptual framework that guides this thesis, current UK policy relating to respiratory health, and the overarching rationale, aims and objectives for this research. This chapter will introduce the data sources used to address my objectives. I will begin by defining administrative data, including the benefits and drawbacks of these types of datasets, before introducing the specific data sources used in my thesis. Next, I focus on an emerging challenge for research that utilises administrative data—patients opting out of sharing their data for research purposes. Throughout this chapter, I will also outline the various research roles I have undertaken as part of my PhD.

3.1. Administrative data

The data used in this thesis comes from administrative datasets, defined here as data that have been generated about citizens through provision and administration of public services. These datasets tend to cover whole populations, since they include all individuals coming in contact with the particular service. They offer information on a large number of people, which is particularly useful for research that compares risk between subgroups or looks at rare events or complex pathways. In some instances, multiple datasets can be linked together, enhancing existing information further. The costs associated with administrative data are typically much lower than research involving de novo data collection, particularly at the same scale, and the time usually spent on data collection is saved. Administrative data may offer a better degree of accuracy than other retrospective designs as the information is normally recorded at or soon after the time of an event. Given that the primary function of these datasets is administration and provision of often universal public services, the risk of non-response bias and loss to follow-up are also minimised.

On the other hand, as these datasets are not designed for the purpose of research, they also present challenges for researchers. Unlike specifically designed cohort studies, researchers generally have no input into data collection and coding practices, and any changes to these over time. Since individuals whose information is contained in administrative databases have not consented to take part in research, a cost and time penalty is associated with applying for
access to these data for research purposes. This is especially the case where individual-level data are required due to concerns about meeting legal requirements for use, confidentiality and avoiding individual disclosure. The datasets are often large, complex and received by researchers in a ‘messy’ format, requiring substantial data cleaning before use.

Data quality may be affected by errors in data collection and, when data linkage is used, when records cannot be linked or are incorrectly linked. These errors can bias the prevalence and distributions of attributes in the dataset, as well as any associational analyses one might perform on the data.

Nonetheless, administrative records provide great opportunities for epidemiological analysis if limitations are recognised and addressed. For this PhD, I used administrative health data from England and Scotland, which I will now describe in further detail.

### 3.2. English Hospital Episode Statistics linked to mortality records

#### 3.2.1. Overview

Hospital Episode Statistics Admitted Patient Care (HES APC) is a database of all inpatient admission records, including day cases, to NHS hospitals in England, and all episodes of care provided in independent hospitals but paid for by the NHS. The data available in HES APC include: clinical information (operations and diagnoses); patient information such as sex, ethnicity and age; administrative information including the dates of admission and discharge; and geographical identifiers such as the Lower Super Output Area (LSOA) of the patient’s address. NHS Digital—the supplier of HES APC data—routinely link admissions to the same patient over time based on NHS number, date of birth, sex, hospital number and postcode.

For secondary uses of the data, pseudonymisation is applied. That is, a unique identifier called the HESID is assigned to each individual and all personally identifying information removed from the dataset (i.e. NHS numbers and full postcodes). HESIDs are unique to each data extract and are available from the financial year 1997/98 onwards.

Births are recorded as hospital admissions in the English NHS, and HES APC captures approximately 97% of births in England that occur in NHS hospitals. There are two separate sources of information on each birth, the delivery episode (maternal record) and the birth episode (baby record), which both contain further details in addition to the main HES APC record. These further details are referred to as the “maternity tail” or “baby tail” and includes information such as birthweight, gestational age, delivery method and birth order (for multiple births). Although mothers and baby pairs are not already linked by NHS Digital when
researchers receive HES APC data, it is possible to apply linkage in the pseudonymised data using a mixture of deterministic and probabilistic methods. Deterministic linkage is where records are linked based on the presence of a set of identical values (e.g. date of birth/delivery, sex, hospital and postcode district). Probabilistic linkage assigns match weights to all possible record pairs based on the probability that a set of variables agree or disagree. Weights above a certain threshold are then selected as matches. Probabilistic methods are essential for linking mother and baby records in HES APC because errors and missing values in the dataset have been shown to lead to a high proportion of missed matches based on deterministic methods alone.

Linking mother and baby records enhances the completeness of the dataset as it is then possible to supplement missing data from one source (e.g. the birth record) with the other (e.g. the delivery record). HES APC allows patients to be tracked via longitudinal hospital records using their HESID. HES APC is routinely linked to the Office for National Statistics (ONS) mortality records, so that the dates and causes of deaths outside NHS hospital settings for people who have a record in HES APC are also available. Alongside the enhancements available through data linkage, the quality of data in HES APC has also improved over time. Burns et al. conducted a systematic review of studies that have assessed the accuracy of hospital episode data across England (56% of the papers), Scotland and Wales. When compared to case or operative notes, the accuracy of primary diagnosis was found to have risen from 73.8% to 96.0% between 2002 and 2009. Since 2004/05, the dataset serves as the basis for a pay-for-performance system for NHS secondary care, which has led to increased reporting of hospital activity and depth of clinical coding. However, for HES APC returns that are not mandated, such as gestational age and birthweight, there remains a high proportion of missing data. Further, consistency of information can vary across and within hospital trusts. For example, 11 of 139 NHS trusts in 2009/10 had relatively poor internal consistency on recording of delivery method.

Another challenge of using HES APC data for research is the proportion of people choosing to opt out of sharing their data for purposes beyond their own care. Where opt-outs are applied, no information is received on these patients, which introduces an unknown amount of bias into the dataset and affects the quality of data linkage. In addition, HES APC does not capture private or charitable hospital activity that is not funded by the NHS. There is no requirement for these hospitals to centrally record activity and, as such, the precise volume and type of care falling under this remit is not freely available. However, since emergency treatment is not typically covered under private medical insurance in the UK and NHS care is free at the point of need, it is expected that very few children are admitted privately for acute
illnesses like bronchiolitis. Therefore, I do not expect this facet to impact on the reliability of findings in this PhD.

3.2.2. Using HES APC in my PhD

Approvals

Approval to use HES APC data was granted by NHS Digital (formally the Health and Social Care Information Centre) for a programme of research within our research unit within the terms of a data sharing agreement, DARS-NIC-393510-D6H1D-v3.2. This agreement is for the use of a de-identified extract of HES APC linked to ONS mortality records for specified objectives. My PhD sits under the second of the three objectives of this work, which is “to determine the risk factors for emergency and recurrent use of secondary care”. As my work fell under current purposes for use, I did not need specific permission from NHS Digital to use these data for my PhD. In addition, as I was using pseudonymised data, my research did not require an ethics review by an NHS Research Ethics Committee.

HES APC birth cohort

When I began my PhD, a birth cohort had already been created from linked HES APC and ONS mortality records by Ania Zylbersztejn. Table 3.1 gives further details of the datasets that comprise the birth cohort, with examples of variables relevant to my PhD. The HES APC birth cohort included linked mother-baby data from 2003, following the introduction of a programme to provide NHS numbers to babies at birth on 29th October 2002, to 2016. Non-English residents (identified using postcode information) were excluded from the dataset. Multiple births were also excluded because of the high possibility of assignment of the same HESID in same sex individuals from multiple births, thereby creating false matches. This is because the algorithm that creates HESIDs relies on sex, postcode and date of birth in situations where an individual’s NHS number is missing.

Table 3.1 Description of datasets comprising the English birth cohort used in my thesis

<table>
<thead>
<tr>
<th>Dataset</th>
<th>Description</th>
<th>Example variables</th>
</tr>
</thead>
<tbody>
<tr>
<td>HES APC</td>
<td>A dataset containing all admissions to NHS-funded hospitals in England. An admission is defined as an activity that requires a hospital bed, including both</td>
<td>HESID, sex, age at admission, admission date, discharge date, diagnoses (up to 20), LSOA of patient address, Index of Multiple</td>
</tr>
</tbody>
</table>
emergency and planned admissions, day
cases, births and deliveries. Deprivation (based on
patient's postcode)

In addition to general admission details, a
tail with 19 variables relevant to the
delivery and labour are included in birth
(baby) and delivery (mother) records.

| ONS mortality records | All deaths registered in England among residents and non-residents. | All causes of death (up to 15), date of death |

I used cohort data for births from 2011 onwards in my analyses and, when I looked at
descriptive data for these years, I noticed a trend towards lower mother-baby linkage rates in
the most recent study years, from 96.4% in 2012/13 to 92.0% in 2016/17. This was despite an
increase in rates of deterministic linkage (i.e. unique links on several characteristics) between
mothers and babies following improvements in hospital recording of data.\textsuperscript{279} Linkage is
particularly important for the geospatial bronchiolitis work in this PhD, as geographical
identifiers are more complete in the mother’s data. I therefore began investigating the potential
source of this unexpected pattern.

**Updating the mother-baby linkage**

I firstly looked at the methodology used to probabilistically link mothers and babies, which is
detailed in Harron et al.\textsuperscript{273} This involves calculating match weights for each record pair based
on the probability that variables across two records randomly match (U-probabilities) and the
probability that there is agreement on a variable given that two records are a true match (M-
probabilities). The match weights had been calculated using U- and M-probabilities derived
from an earlier extract of the data (≤2012) and may not have been as accurate for the newer
more complete data. This led me to redo the linkage for HES years 2013/14 to 2016/17 with
the aim of improving linkage rates. The full methodology is outlined in Appendix 3.1. However,
on completion of this update I still found increasing rates of unlinked babies, as shown in the
‘updated linkage’ column of Table 3.2. Looking again at the raw number of mothers and
babies, I identified that the number of delivery records compared to birth records diverged
significantly from 2014/15 (‘extra birth records’ column, Table 3.2). Even if linkage methods
were perfect, 3.3% of babies born in 2016/17 did not have a delivery record available for
linkage.
Table 3.2. Number of singleton birth records, proportion unlinked to delivery records in original and updated linkage and available delivery records, by financial year

<table>
<thead>
<tr>
<th>Year of birth</th>
<th>Birth records (N)</th>
<th>Initial linkage (%)</th>
<th>Updated linkage (%)</th>
<th>Delivery records (N)</th>
<th>Extra birth records* (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>2012/13</td>
<td>645,937</td>
<td>96.4</td>
<td>96.4</td>
<td>641,080</td>
<td>0.8</td>
</tr>
<tr>
<td>2013/14</td>
<td>626,161</td>
<td>96.0</td>
<td>96.4</td>
<td>623,863</td>
<td>0.4</td>
</tr>
<tr>
<td>2014/15</td>
<td>613,401</td>
<td>92.4</td>
<td>94.0</td>
<td>596,916</td>
<td>2.8</td>
</tr>
<tr>
<td>2015/16</td>
<td>625,065</td>
<td>92.1</td>
<td>93.3</td>
<td>605,643</td>
<td>3.2</td>
</tr>
<tr>
<td>2016/17</td>
<td>617,311</td>
<td>92.0</td>
<td>93.6</td>
<td>597,127</td>
<td>3.3</td>
</tr>
</tbody>
</table>

*Compared to the number of delivery records, calculated as (birth records - delivery records) /delivery records.

Impact of patient opt-outs

This led to my second identified issue with the dataset: patient opt-outs. Since January 2014, patients of the NHS in England have been able to opt out of sharing their patient data for research and planning purposes (previously known as a type 2 opt-out). Accordingly, NHS Digital have applied opt-outs to our HES APC data extract before data release for years 2014/15 onwards. In 2017, when we received the data used in this thesis, the average rate of type 2 opt-outs across England was 2.3%. This is the average proportion of patients we would expect not to appear in our data extract. We do not have details about people who opt out, meaning that it is unclear if pregnant women opt out at the same rate as the general population. However, it is likely that opting out accounts for at least some of the pattern shown in Table 3.2. I discuss the wider impact of patient opt-outs in Section 3.4.

3.3. Scottish administrative health databases

3.3.1. Overview

Health data are organised differently by the NHS in Scotland. Instead of one database with all admitted care, birth records, maternity inpatient cases and other hospital admissions are collected in separate datasets. Patients can be linked across health datasets and over time using their Community Health Index (CHI) number. The CHI number is a unique identification number, equivalent to NHS numbers in England and Wales, that is assigned when a patient first registers with NHS Scotland or at birth. There are several Scottish administrative health databases pertinent to this thesis, which I outline in Table 3.3, below. To enable researchers to connect records belonging to the same patient across these datasets, the Electronic Data Research and Innovation Service (eDRIS) team (part of the Information Services Division of...
the NHS National Services Scotland) assigns a unique, pseudonymised identifier to each individual. For child and maternal datasets, eDRIS provides pre-linked infant and maternal records, with the linkage derived using a combination of deterministic and probabilistic methods. Mother and baby linkage rates reached 98.1% by 2017 in our data extract, emphasising the value of this dataset and the long history of the use of CHI numbers on health records in Scotland.

As noted by Pavis and Morris, “Scotland has some of the best administrative and care data in the world.” Electronic national data from the NHS in Scotland is available for all births from 1981 and, in 2015, less than 2% of Scottish residents used private healthcare. Pavis and Morris write that public engagement has been key in the Scottish model of administrative health research. In response to public attitude and legislative requirements, research datasets are bespoke and project-specific rather than from one permanently linked dataset. Projects are vetted at the local, regional, and national level to ensure there is clear public benefit. Unlike in England, there is no digitalised opt-out service for patients of the Scottish NHS. Instead patients are instructed to opt out by sending “a written request supported by adequate proof of identity” to National Services Scotland—the national NHS Board that provides national strategic support to NHS Scotland.

Vitally to my work on asthma, there is a national individual-level dataset of NHS prescriptions in Scotland that can be linked to other health datasets (see PIS dataset, Table 3.3). This information can be used to identify children who require medication to manage asthma, and was not available for England at the time of data request. In 2009 the quality of the prescription data improved drastically, with reimbursed data capturing 87% of patient identifiers, enabling the majority of items to be linked across other ISD datasets. This figure rose to 96% of reimbursed prescriptions and almost 100% of prescribed and dispensed items by 2014.

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*A note on the changing structure of NHS Scotland.* In 2020, National Services Scotland (NSS) began transitioning to Public Health Scotland (PHS). ISD Scotland, which used to sit under NSS, is now part of PHS and has been renamed “data and intelligence”. I use the former names of these teams/departments throughout this section for consistency.
### Table 3.3. Description of Scottish health datasets

<table>
<thead>
<tr>
<th>Dataset</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>NRS live birth, stillbirth and death registration records</td>
<td>National Records of Scotland (NRS) is the department of the Scottish government that is responsible for vital event registration, the census and national archives. Before April 2011 these records were held at the General Register Office for Scotland. We received birth, death, and stillbirth registrations in separate files.</td>
</tr>
<tr>
<td>CHI registry (emigration records)</td>
<td>CHI status (no link to CHI, emigrants or untraced, death or currently on CHI) and date of emigration are recorded in a CHI register, which is held by NHS Scotland. This information is derived from GP records.</td>
</tr>
<tr>
<td>SBR and SMR11 birth records</td>
<td>Scottish Birth Records (SBR) are a singular electronic record capturing stillbirths and neonatal care received by babies in Scotland, including home births. It does not include diagnostic information. This dataset replaced the old birth record dataset (SMR11) in 2002/03.</td>
</tr>
<tr>
<td>SMR02 delivery records</td>
<td>SMR02 is submitted to ISD Scotland by maternity hospitals and contains inpatient and day case activity relating to obstetrics. Information about the mother, deliveries and new-born babies is available in this dataset.</td>
</tr>
<tr>
<td>SMR01 hospital admissions</td>
<td>SMR01 includes all inpatients and day cases discharged from hospitals in Scotland (apart from obstetric wards, SMR02, and psychiatric wards). The records include diagnostic, procedure and operation information, as well as non-clinic information such as location and length of stay.</td>
</tr>
<tr>
<td>PIS</td>
<td>The Scottish National Prescribing Information System (PIS) is a dataset of NHS medicines prescribed and dispensed within the community setting in Scotland. Prescriptions dispensed within hospitals are not included. Information includes date and place of prescription and dispensing activity, and drug information including British National Formulary (BNF) codes.</td>
</tr>
</tbody>
</table>
3.3.2. Using Scottish Administrative Health Databases in my thesis

Approvals

Applying for Scottish healthcare data involved submitting a Public Benefit and Privacy Panel for Health and Social Care application form to National Services Scotland (now under Public Health Scotland). My primary supervisor, Pia Hardelid, led this process with input on the content from myself and another colleague. I also contributed to this process by designing a questionnaire to seek the perspectives of patient interest groups specifically on the topic of linking data across databases. We sent this questionnaire to the Great Ormond Street Hospital parent carer advisory group and a stakeholder who replied to an advertisement in Asthma UK’s monthly bulletin (the questionnaire is included in Appendix 3.2). The responses revealed a primarily positive view of data linkage for the purposes outlined but raised questions about the lack of wider dissemination of such methods. We received full Public Benefit and Privacy Panel for Health and Social Care approval on 21st November 2018, and the full data extract on 11th December 2019.

Updating the Scottish birth cohort

A birth cohort, including births from 1981 to 2015, had previously been created from Scottish health data by a colleague, Maximiliane Verfürden.\textsuperscript{296,297} I adapted the code created by Maximiliane and other colleagues to clean the new files and create an updated birth cohort including births from 1981 to 2017. I used the NRS birth registration data as the core birth file and added information from other datasets to build the birth cohort as shown in Figure 3.1. In addition, I cleaned and structured the PIS dataset (which had not previously been held by our team), so that information about medication can be easily linked with mothers and children in the main cohort via their mother-ID or child-ID, when needed.

There were 2,203,042 NRS birth registrations in total over the 36-year period. I excluded 12,951 (0.6%) records with an unknown child-ID in the NRS birth registration dataset, as these records cannot be linked across datasets. The number of unknown child-IDs decreased with year of birth and the vast majority (84%) were from births before the year 2000. I excluded 33 stillbirths, identified using the “outcome of pregnancy” variable in SMR02, and 332 infants registered with a non-Scottish address at birth, identified through postcode area. Lastly, 538 duplicate records were excluded from the cohort. I changed improbable values to missing where: maternal age outside of 10 and 60 years (n=14); gestational age was outside of 22 and 45 weeks (n=178); birthweight was less than 100 grams (n=100); and date of death occurred before date of birth (n=14). The resultant whole country birth cohort of 2,189,188
infants covered 99.1% of all births in Scotland over this period when compared with official vital event statistics.\textsuperscript{298}

### Master file

#### NRS birth registrations, 1981 to 2017
- Carstairs index of deprivation 1981 to 2011*  
- Child-ID  
- Birth date  
- Mother country of residence  
- Postcode area*  
- Sex*  
- SIMD 2004 to 2012\textsuperscript{†}  

*supplemented with information from other datasets where missing; †SIMD = Scottish index of Multiple Deprivation

#### Birth records
- SMR11, 1981 to 2002/03  
  - Admission date  
  - Birthweight  
  - Carstairs index 1981-2011  
  - Child-ID  
  - Diagnoses (up to 10)  
  - Discharge date  
  - Ethnicity group  
  - Intensive care unit stay  
  - Sex  
  - SIMD 2004-2012

#### Death records
- NRS death registrations, 1981 to 2018  
  - Child-ID  
  - Death date  
  - Primary cause of death  
  - Secondary causes of death (up to 10)

#### Delivery records
- SMR02, 1981 to 2017  
  - Apgar score  
  - Birthweight  
  - Carstairs index 1981-2011  
  - Delivery date  
  - Delivery mode  
  - Ethnicity group  
  - Gestational age  
  - Hospital  
  - Maternal age  
  - Mother-ID  
  - Number of births  
  - Outcome of pregnancy  
  - Parity  
  - SIMD 2004-2012  
  - Sex of baby  
  - Smoking status

#### Emigration records
- Emigration records, 1981 to 2018  
  - CHI status  
  - Emigration date

#### Hospital admissions
- SMR01, 1981 to 2018  
  - Admission date  
  - Discharge date  
  - Main condition  
  - Other conditions (up to 5)

---

**Figure 3.1.** Scottish health datasets and relevant variables used to create birth cohort
3.4. **National data opt-out programme: consequences for maternity statistics in England**

Whilst exploring issues in the HES APC birth cohort, I became interested in understanding the wider potential consequences of patient opt-outs. I noted that the existing discussion of the English NHS patient data opt-out programme mostly neglected a positive view of why sharing data may be for the benefit of population health. I therefore decided to write a commentary article exploring the impact of NHS England patient opt-outs on maternity service statistics. I sought feedback on an early draft of the paper from members of a patient and public working group based at the UCL Institute of Health Informatics (formerly the Farr Institute, see Appendix 3.3). Ten members of the group gave comments that helped to shape the arguments below. The full article, which was published in the International Journal of Population Data Science, is available in Appendix 3.4.

3.4.1. Background

NHS Digital, the data capture organisation within the English NHS, collects and stores some of the information recorded when individuals receive health or adult social care in England. The data are used for the purpose of improving both individual care, through improved record sharing across health providers, and the wider population’s health and care, through activities such as planning services, monitoring epidemics and researching health conditions and treatments.\(^{272}\) As part of the population-based work, NHS Digital oversees the strictly controlled release of patient information to third parties including NHS providers and commissioners, university researchers, charities and companies that are partnered with the NHS. These organisations go through a lengthy application process to gain access to the data and follow stringent protocols when storing and analysing the data. Further, personal identifiers, such as names and NHS numbers, are removed in all circumstances apart from where specific patient consent is given or where required by law.\(^{272}\)

The quality and completeness of English administrative health data, and the quantity of health research based on these data, have been increasing over time.\(^{299}\) However, a new barrier to the quality of administrative data is emerging in the form of opt outs. That is, patients choosing not to share their information beyond the NHS for anything other than their direct care and treatment. Opting out of NHS data sharing was first made available for patients of the English NHS in January 2014 following recommendations in Dame Fiona Caldicott’s 2013 information governance review.\(^{300,301}\) In May 2018, a new online consent platform called the National Data Opt-Out service was launched by the NHS. This platform was intended to be simpler and
easier to access than the previous model; however, based on low numbers of uptake four months after its launch, it has been argued that not enough patients are aware of the scheme.\textsuperscript{302}

As at 1 December 2018, the average national data opt out rate across England was 2.8\%.\textsuperscript{303} This reasonably low proportion of missing records would be less concerning if the characteristics of those opting out were the same as those not opting out (i.e. missing at random). However, top-level demographic information published by NHS Digital shows that opting out is not uniformly distributed by age, sex or geographical area.\textsuperscript{303} Figure 3.2 presents opt out rates by Clinical Commissioning Group (CCG), showing clear variation across the country. Notably, twelve CCGs had opt out rates higher than 5\% and one CCG had a rate of 10.1\%. At the GP practice level, there were instances where the entire patient population have opted out, which raises significant questions about whether the patients in these practices explicitly opted out for themselves.\textsuperscript{301,303}

![Figure 3.2. Rates of patients opting out, by CCG: England (with inlay map of London CCGs), as at December 2018. Data from NHS Digital.\textsuperscript{40}](image)
The 2016 Caldicott review (the third in the series) stated that ‘patients have a right under the NHS Constitution to request that their personal confidential information is not used beyond their direct care’.\textsuperscript{304} I am not disagreeing with this principle. Legitimate concerns about the security of patient data have been voiced by both the general public and experts in the medical field, particularly in response to the contentious (and discontinued) care.data scheme.\textsuperscript{305,306} However, in parallel with information about this choice, patients should have complete and transparent information about the uses and potential advantages of sharing their information. Using a simple simulation based on publicly available data, the aim of this research is to exemplify the potential impacts of patient opt outs on maternity information and research.

3.4.2. Application to NHS maternity statistics

As introduced in Section 3.2, HES APC is the most comprehensive source of information on all births and deliveries in England.\textsuperscript{274} Delivery records from this dataset are used for many purposes, including to create resources for parents-to-be, to evaluate and improve maternity care provision and to investigate multiple risk factors for ill-health in mothers and babies.\textsuperscript{307} Tools, such as the Birth Choices tool from Which?,\textsuperscript{308} have been created utilising this data to aid expectant parents when making maternity choices. The National Maternity and Perinatal Audit, which was set up in 2016 to evaluate quality in NHS maternity services, uses maternity data from HES linked with data from each maternity unit to provide a range of statistics comparing outcomes at maternity unit level.\textsuperscript{309} In addition, HES APC has been used for maternal and child health research to examine, for example, factors explaining excess child mortality in England and the safety of surgical procedures during pregnancy.\textsuperscript{284,310} Individual level data are necessary for this type of research, since this allows multiple risk factors for maternal and child outcomes to be taken into account.

To exemplify the impact of biased data on maternal information, I calculated the potential rates of two health indicators had opt outs not been applied to HES delivery data. I downloaded publicly available CCG-level information from Public Health England’s Fingertips Child and Maternal Health Profiles for one common and one rare outcome: the proportion of deliveries with caesarean sections which occur in approximately 27% of births nationwide; and births with very low birth weight (<1500g), which has a nationwide prevalence of 1.2%.\textsuperscript{311} I chose three CCGs for the example, each with a different rate of opt outs as published at 1\textsuperscript{st} December 2018 by NHS Digital:\textsuperscript{312} Bradford City with a 0.3% opt out rate; Merton with a 2.8% opt out rate; and Oldham with a 10.1% opt out rate. Under the assumption that the women in the maternity dataset opted out at the same rate as the whole CCG population, I modelled three scenarios based on the rate of events in the women that opted out: 1) no events; 2) events occurring at the same rate as women who did not opt out; 3) all had an event. Mirroring
methods used by Public Health England, \(^{313}\) 95% confidence intervals were calculated in Stata 15.0\(^{314}\) using the Wilson Score method. Microsoft Excel 2013 was used to create graphs.

Table 3.4 displays published and simulated event rates for caesarean sections and infants born with very low birth weight, by CCG. Where patients opting out have the same rate of events as patients not opting out (scenario 2, Table 3.4), published and simulated event rates do not differ. However, the range of potential rates between the extremes of scenario 1 (no new events among women opting out) and scenario 3 (all women opting out had an event) shown in Figure 3.3 highlights the huge potential for misleading information where the frequency of events differs between patients that do and do not opt out. These scenarios show that both the CCGs with average (Merton) and high (Oldham) rates of opting out could be presenting misleading information. For births with very low birth weight (the rarer event), the rate of births with this outcome could potentially be 9-fold higher than the published rate in Oldham. The non-random nature of opt-outs, therefore, has potentially large implications for the outcomes of public health information and monitoring. At an NHS trust level, information bias could produce flawed outputs in audits intending to suggest improvements and highlighting good practice in maternal care and infant outcomes. These biases will also affect the reliability of findings on maternal and children’s health research. As shown in this simple simulation, this is particularly the case where less common (but often more serious) outcomes are studied.

Problematically, the bias introduced into datasets with opt-outs applied cannot be treated with the same statistical methods used to treat missing data. Multiple imputation, a method which is commonly applied to deal with missing data, relies on the assumption that the missing information can be explained by differences in the observed data.\(^{315}\) However, once patients have opted out, their data is completely removed from the dataset (i.e. complete case removal) meaning that researchers cannot account for systematic differences. Using multiple imputation in these instances may actually add further bias to results.\(^{315}\) Methods used in population-based surveys to overcome biases of non-consent, such as weighting adjustments or simulation studies ideally require detailed data on the population (e.g. by gender, age, deprivation level and local area) who have opted out so that correct weights can be derived—information currently not published by NHS Digital. Research to determine whether those who have opted out of sharing data are different in terms of socio-demographic and health characteristics, in specific populations such as expectant mothers, would help in applying such methods to tackle data missing due to opt outs.
Table 3.4. Published and simulated number of events, patients and event rates, by indicator and CCG

<table>
<thead>
<tr>
<th>Indicator</th>
<th>CCG</th>
<th>Published maternity data</th>
<th>Opt out rate (95% CI)</th>
<th>New event rate (95% CI)*</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Events</td>
<td>Deliveries</td>
<td>Event rate (95% CI)</td>
</tr>
<tr>
<td>Very low birth weight</td>
<td>Bradford City</td>
<td>36</td>
<td>1,657</td>
<td>2.17 (1.57, 2.99)</td>
</tr>
<tr>
<td></td>
<td>Merton</td>
<td>39</td>
<td>3,219</td>
<td>1.21 (0.89, 1.65)</td>
</tr>
<tr>
<td></td>
<td>Oldham</td>
<td>39</td>
<td>3,195</td>
<td>1.22 (0.89, 1.66)</td>
</tr>
<tr>
<td>Caesarean section</td>
<td>Bradford City</td>
<td>478</td>
<td>2,042</td>
<td>23.6 (21.8, 25.5)</td>
</tr>
<tr>
<td></td>
<td>Merton</td>
<td>948</td>
<td>3,265</td>
<td>29.0 (27.5, 30.6)</td>
</tr>
<tr>
<td></td>
<td>Oldham</td>
<td>792</td>
<td>2,797</td>
<td>28.3 (26.7, 30.0)</td>
</tr>
</tbody>
</table>

*Scenario 1 (no new events) = published events/(deliveries + extra deliveries) where extra deliveries = published deliveries/(1-opt out rate)-published deliveries; scenario 2 (average events) = ((extra deliveries*event rate)+published events)/(deliveries + extra deliveries); scenario 3 (max. events) = (events+ extra deliveries)/(deliveries + extra deliveries)

Figure 3.3. Observed and simulated rates of births with very low birth weight (<1500g, 2016/17) and deliveries by caesarean section (2016): by CCG
3.4.3. Are opt outs inevitable?

Public health research in the Nordic countries demonstrates that data sharing can be achieved with buy-in from citizens and at great value to clinical research. In these countries, residents are assigned a personal identity number from birth, which can be used to track individuals across time and generations. Unlike the UK equivalent (NHS number), which is used only in health and social care settings, personal identity numbers are used across a multitude of sectors. This means that individual data can be linked across health, education and social security datasets, for example, providing high quality comprehensive information on risk factors for ill-health and other outcomes. The safeguards in place are very similar to those in England. In contrast, however, there is broad public general awareness and acceptance of the use of individual data in research and a long-standing culture of trust in public services and data donation for the good of the population.

There is some evidence from England to suggest that greater knowledge of research processes and safeguards improves the likelihood of acceptance of electronic health records for research being used without explicit consent. An electronic real-time dataset integrating primary and secondary care was successfully implemented over a decade ago in Salford, Manchester. All patients were sent a letter with information about the project and whether they would like to opt out. Reportedly, less than 0.2% of the nearly quarter of a million patients chose to opt out. In contrast, information about the now withdrawn care.data scheme to integrate primary and secondary care records was disseminated by generic leaflets, which were reportedly not seen by the majority of the population. Some additional strategies were used to provide information about the national opt out scheme to the public. A national radio campaign ran for 6-weeks after the launch of the new system and NHS Digital's website now links to Understanding Patient Data, a website providing “objective information about how patient data is used” that is run by Wellcome Trust. However, given that some of the opt-outs may have been driven by GP practice rather than patient-level decisions, it is not clear to what extent whether it was the NHS information campaign directed at patients (or indeed the absence of it) that led to the 2.8% opt-out rate, or a lack of buy-in from clinicians.

What is known is that when patients choose to opt out of sharing data beyond their direct care, the reliability of service information and evaluation and wider research based on electronic health records is diminished. NHS Digital is beginning to advertise the benefits of sharing NHS data for research and planning purposes, but there are ways this could be improved. NHS Digital could start by listing examples of how health care data have been used for research, as is available in other NHS held datasets (e.g. CPRD). NHS Digital currently publishes a data release register, but not a data output register. Other examples of dissemination
include the University of Manchester’s citizen’s jury on health records and an animation created as part of the #datasaveslives campaign by the Farr Institute. However, the impact and wider reach of these schemes are not clear and, arguably, information about the benefit of data sharing can only go so far in raising public confidence.

Evidence from the research literature and reflected in media coverage suggests that there is unease about the potential of commercial entities, such as pharmaceutical and insurance companies, making profit from NHS data. This is in contrast to the largely positive view of university researchers or NHS staff making use of this data. Clear examples of ‘acceptable and unacceptable purposes’ for which data can and cannot be used, as called for by the Wellcome Trust could help to reassure the public. Further, clinicians play a pivotal role in the discourse of patient consent to use NHS data for research. Research to better understand these concerns and how they could be addressed could be an important step to improving information campaigns.

Overall, there needs to be greater transparency through clear and detailed information on: who can apply to use NHS data and for what reasons; the safeguards in place to protect individual information; and, importantly, the wider consequences of opting out on population health research and public health service information. Only with this information can individuals be expected to make an informed decision about opting out of sharing their data.

3.5. Chapter summary

In this chapter, I have outlined the composition of the English and Scottish birth cohorts that will be used to conduct research throughout this thesis. The English birth cohort is a pre-existing dataset derived from HES APC data linked to ONS mortality records. I contributed to this dataset by updating the linkage between maternal and baby records, which led to an increase in the data completeness of the cohort. The Scottish birth cohort, which I updated from an older version of the dataset, is composed of several health databases including national birth and death registrations, hospital admissions and delivery records. The final part of this chapter focuses on the emergent issue of data opt outs and the potential implication of this on maternity research and information.
Chapter 4. Geospatial and seasonal variation of bronchiolitis in England

Recap and chapter overview

Chapters 1 to 3 presented an outline of the research literature and an introduction to the datasets in this PhD. This chapter is the first of two that focus on the seasonality of bronchiolitis admissions. Using a birth cohort created from the English hospital data introduced in Chapter 3, I explore spatial variation in the seasonality of hospital admissions for bronchiolitis and its association with local demographic characteristics. This is the first step in answering thesis objective 1:

*To describe the association between seasonality, early life socioeconomic position and bronchiolitis admission rates among infants across England.*

An article based on the contents of this chapter was published in Thorax (see Appendix 4.1) and the Stata code for the analyses is published online at [https://github.com/UCL-CHIG](https://github.com/UCL-CHIG).

### 4.1. Background

Geographical location is a prominent source of variation in rates of hospital admissions for bronchiolitis in England.\(^4,15,45\) On average, between 1999 and 2011, a 5.3-fold difference in the rate of hospital admissions for bronchiolitis among infants was reported across the 352 Local Government Areas of England.\(^4\) Another study that looked at admissions across the 152 Primary Care Trusts in England (the local NHS administrative bodies that were abolished and replaced by Clinical Commissioning Groups, CCGs, in 2013) between 2007/08 and 2010/11, described a 15-fold difference in the rates of admissions for bronchiolitis among children under 2 years of age.\(^327\) An overarching factor proposed to explain this extreme geographical variation lies with socioeconomic differences in the composition of the population at the area-level. However, these studies noted only modest associations between higher socioeconomic deprivation (measured by the index of multiple deprivation) and bronchiolitis rates; \(r^2=0.24\) at the Local Government Area-level and \(r^2=0.33\) at the Primary Care Trust-level.\(^4,327\) Instead, the authors of these papers write that it is likely that clinical factors, particularly admission thresholds, are the main driver of the variation in admission rates noted.
However, these geographical studies did not consider the impact of seasonality when assessing spatial variation in admissions for bronchiolitis across England. Seasonality, defined as cyclic changes in disease occurrence, is a central component to the patterns of bronchiolitis admissions observed in England. In temperate maritime climates such as in the UK, respiratory syncytial virus (RSV, the dominant pathogen causing bronchiolitis) epidemics occur during the colder months, with a parallel trend in hospital admissions for bronchiolitis.

By including sine and cosine functions in regression models of times series data (analysis known as harmonic regression), generic seasonal patterns can be removed from estimates and underlying differences in trends detected. Harmonic regression analysis also allows for the calculation of additional area-specific disease components, such as the timing of the highest disease rates and the epidemic duration. Researchers in Australia and the USA have used this type of analysis to look at population factors associated with features of RSV epidemics among children. For example, Noveroske et al. found that earlier epidemic timing was associated with areas of Connecticut, USA, that were more densely populated and had a higher proportion of black people in the population (used in this study as an indicator of socioeconomic deprivation). This type of analysis, yet to be conducted in a UK setting, can be used to plan interventions, such as whether differential timing of palivizumab prophylaxis based on area or population characteristics is necessary.

The aim of this Chapter is to describe the spatial variation in the seasonality of hospital admissions for bronchiolitis across England.

There are two objectives for the Chapter:

**Objective 4A** To demonstrate how seasonality of bronchiolitis admissions amongst infants varies across region and CCG in England.

**Objective 4B** To assess how epidemic timing and rates of admissions (after accounting for seasonal differences) are associated with population characteristics between CCGs.

### 4.2. Methods

#### 4.2.1. Data sources and study cohort

This study uses hospital episode statistics admitted patient care (HES APC) data linked to Office for National Statistics (ONS) mortality records as introduced in Chapter 3. HES APC is a database of all hospital inpatient admissions funded by the English NHS and captures approximately 97% of all births in England. NHS Digital, the body that houses and supplies
access to this database, links patient records over time and provides each database extract with a pseudo-anonymised identifier unique to each individual (called a HESID), enabling researchers to create longitudinal patient cohorts. I also used CCG-level data, namely index of multiple deprivation (IMD) scores and population size, which I downloaded from the Ministry of Housing, Communities and Local Government (MHCLG) and ONS websites, respectively.\textsuperscript{333,334}

A birth cohort of children born from 1st January 2011 to 31st December 2016 was derived from the dataset previously created by my colleague, Ania Zylbersztejn.\textsuperscript{284} This dataset did not contain infants from multiple births or stillbirths. Using the linkage methods developed by Harron et al.,\textsuperscript{273} I successfully linked 95.0\% of birth records to maternal delivery records (see Appendix 3.1 for further details). This enabled me to complete missing geographical information for infants as recorded at birth/delivery. Non-English resident children were excluded since they cannot be followed up in English hospital data. Our dataset does not contain postcodes or addresses, so non-residency was identified using lower super output area (LSOA). LSOAs not beginning with “E”, which signifies an English LSOA, were used to define non-resident status. Infants with missing geographical data were also excluded from this study.

Children were followed from birth or 1\textsuperscript{st} January 2012, whichever was later, to their 1\textsuperscript{st} birthday, date of death, (estimated) date of emigration or 31\textsuperscript{st} December 2016, whichever came first. This study period ensured that the study had follow-up for children aged up to 12 months in all study years, including the first (2012). Emigration was defined as the presence of a non-English LSOA in any admission during follow up. The emigration date was set as the mid-point between the child’s date of birth and the date at which the non-resident admission occurred. I focused on admissions in the first year of life since more than 90\% of bronchiolitis admissions occur in the first year of life and two thirds of admissions are in children less than 6 months old.\textsuperscript{6,261}

4.2.2. Study outcome

My study outcome was the number of hospital admissions for bronchiolitis, recorded as either the primary or one of the 19 secondary diagnoses in HES APC. Diagnoses in HES APC are coded using the International Classification of Diseases, version 10 (ICD-10), and I identified bronchiolitis admissions using ICD-10 J21 codes indicating acute bronchiolitis. All J21 subcategories were included in this definition: J21.0 acute bronchiolitis due to RSV, J21.1 acute bronchiolitis due to human metapneumovirus, J21.8 acute bronchiolitis due to other specified organisms and J21.9 acute bronchiolitis, unspecified. In this study, sequential
admissions for bronchiolitis by the same child were assumed to be associated with the same infection if the start date of a new admission was within 14 days of the previous admission’s discharge date. In these instances, only the first of these admissions were included in the number of admissions.

Identifying RSV-related bronchiolitis using HES APC data

HES APC data does not provide reliable information about the aetiology of bronchiolitis cases. As mentioned, there are several sub-categories of J21 available, including one that specifies that the cause of the condition is RSV. However, specific bronchiolitis diagnoses are poorly coded in HES APC because laboratory testing is not routine for children presenting to hospital in England with bronchiolitis.45,335 This follows advice from the National Institute for Health and Care Excellence (NICE), who recommend against the routine testing of viral aetiology in their 2015 viral bronchiolitis guidelines for clinicians in England.3 Similar guidance by the American Academy of Pediatrics provides a rationale for this recommendation: “at the individual patient level, the value of identifying a specific viral aetiology causing bronchiolitis has not been demonstrated”.336 p.e1480 However, other methods have been used to ascertain the proportion of bronchiolitis cases in infants that are attributable to RSV. Reeves et al. jointly modelled of HES APC and laboratory surveillance data over a 5-year period from 2007 and estimated that 82% (95% CI 79% to 87%) of admissions with a primary diagnoses of bronchiolitis among children under 6 months and 70% (95% CI 66% to 75%) in children between 6 and 12 months of age could be attributed to RSV.6

4.2.3. Covariates

Year and week of the year at admission were derived from each bronchiolitis admission record. All other covariates were based on each infant’s recorded residence at birth.

Government region was used to describe broad patterns across England and CCGs to examine local areas. There are nine government office regions in England, created to delegate functions from national government in 1994, but now used only for administrative and statistical purposes.337 In ascending order of population size these are the North East, East Midlands, Yorkshire and the Humber, South West, West Midland, East of England, North West, London, and the South East. CCGs, introduced in England in 2013, are NHS bodies responsible for planning and commissioning for local areas in England.333 There were 209 CCGs in England over the study period with mid-2016 populations ranging from 68,187 in Corby (Northamptonshire) CCG to 898,025 in Northern, Eastern and Western Devon CCG.
The number of CCGs within each government office region ranged from 11 CCGs in the North East to 39 CCGs in the South East.

I calculated annual population density estimates—the number of residents per square kilometre—for each CCG by dividing annual CCG population sizes by the area of each CCG in square kilometres (estimated using polygon areas in QGis).\textsuperscript{333} The average population size over the five-year study period was assigned to each CCG. IMD is an overall measure of deprivation experienced by people within a neighbourhood, with a higher score indicating higher levels of deprivation.\textsuperscript{65} The composite score is constructed by combining weighted scores from seven domains of deprivation; income, health and disability, education, skills and training, barriers to housing and services, crime and living environment, with measures mainly derived from the 2011 Census. I used IMD scores for 2015 in this study, summarised across CCGs. These summaries are calculated by averaging the population-weighted IMD scores of the LSOAs contained within a CCG (and are available to download directly from GOV.UK).\textsuperscript{334,338}

### 4.2.4. Statistical methods

Stata 15.0\textsuperscript{314} was used for data analysis. QGIS 2.18,\textsuperscript{339} a free open-source geographic information system, was used to present geospatial data and Microsoft Excel 2013 was used to create graphs. The Stata code for the harmonic regression analysis can be found in Appendix 4.2.

Weekly incidence rates of bronchiolitis admissions for each study year, by region and CCG, were calculated by dividing the number of admissions by person-time at risk. Rates are expressed as admission-based rates per 1000 infant-years.

**Objective 4A: Seasonality of bronchiolitis admissions across region and CCG**

To explore geographical variation at the regional level I fitted a fixed effects Poisson regression model to the regional count data. I used Poisson regression rather than negative binomial regression because there was no evidence of over-dispersion. I used robust standard errors to protect against model misspecification. The impact of seasonality on rates of admissions at the regional level were modelled using a harmonic function of time in weeks ($t$) expressed as: $\beta_1(\sin(2\pi t/T)) + \beta_2(\cos(2\pi t/T))$, where $T =$ number of periods within one cycle (i.e. 1 year = 52.14 weeks).\textsuperscript{340,341} I adjusted for year of admission to account for the annual increasing trend in bronchiolitis admission rates.\textsuperscript{4} I included interaction terms between region and the sine and cosine functions in the region-specific analyses to allow for effect modification of seasonality by region, formally tested via a Wald $\chi^2$ test.
Using the estimated harmonic function coefficients, I calculated the following quantities for each region’s epidemic curve at reference values of the other covariates:

- the amplitude (log) \( \hat{\gamma}_j = \sqrt{(\hat{\beta}_1 + \hat{\delta}_1)^2 + (\hat{\beta}_2 + \hat{\delta}_2)^2} \);
- the phase (in radians) \( \hat{\psi}_j = \arctan(\hat{\beta}_1 + \hat{\delta}_1) / (\hat{\beta}_2 + \hat{\delta}_2) \);
- and the peak week \( \hat{P}_j = 52.14 \times \left( \frac{\hat{\psi}_j}{2\pi} \right) + 1 \)

For descriptive purposes, I also calculated the epidemic duration as the number of weeks between the first of three consecutive weeks with increasing predicted rates and the first of three consecutive weeks with decreasing predicted rates. See Box 4.1 for full model parameterisation.

As outlined in Box 4.1, there were two parts to the CCG-level analyses. In stage 1, I fitted a multilevel (i.e. mixed-effects) Poisson model to the individual admission records. I included the harmonic functions as specified above, and allowed for clustering at the CCG level via random effects for the intercept and the harmonic function parameters. Using the estimated harmonic function coefficients, I calculated CCG-specific average annual peak week as \( \hat{P}_i = 52.14 \times \left( \frac{\arctan((\hat{\beta}_1 + \hat{\eta}_1)/(\hat{\beta}_2 + \hat{\eta}_2))/2\pi}{2\pi} \right) + 1 \), where \( i \) indicates CCG. The CCG-specific departures from the model intercept (i.e. the random effects for the intercept, \( \eta_{0i} \)) across England were exponentiated to obtain CCG-specific incidence rate ratios (IRRs) relative to the population mean.

**Objective 4B: Association between CCG-level epidemic timing, rates of admissions and demographic characteristics**

In stage 2, with CCGs treated as units of analysis, multivariable linear regression models were fitted to estimate the mutually adjusted association of population density and deprivation score with, separately, the CCG-specific peak week and IRRs of bronchiolitis admissions predicted in stage 1. Population density was highly positively skewed and therefore first log-transformed to reduce the impact of very high values. I added quadratic terms to allow for non-linear associations with the explanatory variables and included interactions between them, comparing model specifications via likelihood ratio tests. Alongside regression coefficients, I report the proportion of total variance in each dependent variable explained by the model using eta-squared (\( \eta^2 \)) and the relative contribution of each model component using partial eta-squared (\( \eta_p^2 \)). To assess the impact of ignoring the uncertainty of the predicted values from the first step I computed bootstrapped standard errors from 1,000 bootstraps (from the two
stages of the analysis). I calculated bootstrapped standard errors using a 20% random sample of the data because of computational constraints due to the size of the complete data.

Sensitivity analyses

These models assumed no spatial dependency of the CCG-level residuals. To assess this assumption I calculated Moran’s I, which is a measure of how related the values of a variable are based on the locations where they were measured (with -1 indicating perfect dispersion, 0 no autocorrelation and 1 perfect clustering). Due to the large difference in admission rates within London compared with the rest of the country, sensitivity analyses were conducted to assess the robustness of the results when London-based CCGs were excluded from analyses. To assess the impact of repeated events by the same child, I replicated the analyses including only the first admission per child.

Box 4.1. Models’ specifications and derived parameters

Region-specific analyses

Let \( n_{jk}(t) \), the number of bronchiolitis admissions in region \( j \) and year \( k \) observed at time \( t \) (measured in weeks), follow a Poisson distribution with rate \( \lambda_{jk}(t) = E(n_{jk}(t))/N_{jk}(t) \) where \( N_{jk}(t) \) denote the person-time at risk in region \( j \) and in year \( k \) at time \( t \). I modelled this rate, after log-transformation, as a function of region, year and time of admission as follows:

\[
\log(\lambda_{jk}(t)) = \beta_0 + \beta_1 \sin(2\pi t/T) + \beta_2 \cos(2\pi t/T) + \sum_{j=1}^{9} \alpha_j I_{\text{region}=j} + \sum_{j=1}^{9} \delta_{1j} \sin(2\pi t/T) I_{\text{region}=j} + \sum_{j=1}^{9} \delta_{2j} \cos(2\pi t/T) I_{\text{region}=j} + \sum_{k=2012}^{2016} \theta_k I_{\text{year}=k}
\]

Where: \( T \) is the length of period within one harmonic cycle (i.e. 1 year = 52.14 weeks); and \( I_{x=x} \) is the binary indicator of the variable \( X \) taking value \( x \). The parameter \( \beta_0 \) is the intercept, \( \beta_1 \) and \( \beta_2 \) are harmonic function coefficients, \( \delta_{1j} \) and \( \delta_{2j} \) are region-specific harmonic function coefficients, and the parameters \( \alpha_1, \delta_{11}, \delta_{21} \) and \( \theta_1 \) are all constrained to be zero to deal with the collinearity of the binary indicators.

CCG-specific analyses: stage 1
Let the number of bronchiolitis admissions in CCG \( i \) and year \( k \), observed at time \( t \) (in weeks), \( n_{ik}(t) \), follow a Poisson distribution with rate \( \lambda_{ik}(t) = E(n_{ik}(t))/N_{ik}(t) \), with \( N_{ik}(t) \) denoting the relevant person-time at risk. Given the large number of CCGs, the following multilevel model with random intercepts and random harmonic coefficients was specified for the log rate:

\[
\log(\lambda_{ik}(t)) = (\beta_0 + \eta_{0i}) + (\beta_1 + \eta_{1i}) \sin(2\pi t/T) + (\beta_2 + \eta_{2i}) \cos(2\pi t/T) + \sum_{k=2012}^{2016} \theta_k I_{year=k}
\]

Where: \( \eta_{0i} \) = random effect component of the intercept; \( \eta_{1i} \) = random effect component of the sine parameter; \( \eta_{2i} \) = random effect component of the cosine parameter; and \( \theta_1 = 0 \) as before. The model's predicted CCG-specific parameters were then used to derive CCG-specific peak time, \( \hat{p}_i \), and CCG-specific amplitude, \( \hat{g}_i \).

**CCG-specific RSV analyses: stage 2**

These predicted values were then related to predictors in two separate linear regression models:

\[
\hat{p}_i = \alpha_{10} + \alpha_{11} z_i + \alpha_{12} w_i + \epsilon_{1i} \quad \hat{g}_i = \alpha_{20} + \alpha_{21} z_i + \alpha_{22} w_i + \epsilon_{2i}
\]

Where: \( \alpha_{10}, \alpha_{20} = \) intercepts; \( \alpha_{11}, \alpha_{21} = \) IMD coefficients; \( \alpha_{12}, \alpha_{22} = \) population density coefficients; \( z_i = \) CCG-specific IMD; \( w_i = \) CCG-specific log-population density; and \( \epsilon_{1i}, \epsilon_{2i} = \) random error terms, assumed to be independently distributed.

### 4.3. Results

#### 4.3.1. Descriptive statistics

Figure 4.1 shows the derivation of the final study cohort. Of the 3,808,247 singleton children born between 2011 and 2016, 81,234 (2.1%) were excluded from the final study cohort. In total, 55,801 (1.5%) infants had missing data for geographical information (region and/or CCG), and the odds of missingness decreased by year of birth (OR 0.92, 95% CI 0.91 to 0.92). The final cohort consisted of 3,727,013 children, of which 34.4% had a London or South East address recorded at the time of birth (Table 4.1) and the average follow-up time per infant was 304 days. Between 1 January 2012 and 31 December 2016, 3.7% of infants (n=139,532) had at least one admission to hospital for bronchiolitis. There were 155,485 admissions for
bronchiolitis by cohort members in total, an average annual admission-based rate of 50.1 per 1000 infant-years (95% CI 49.9 to 50.4).

Bronchiolitis admission rates increased from 46.9 per 1000 infant-years (95% CI 46.4 to 47.4) in 2012 to 58.4 per 1000 (95% CI 57.8 to 59.1) in 2016. This was equivalent to a 5.2% annual increase of admissions, from 29,853 to 36,028, between 2012 and 2016. Annual admission rates were highest in the North West (68.7 per 1000 infant-years, 95% CI 67.9 to 69.5) and North East (63.8 per 1000 infant-years, 95% CI 62.5 to 65.1); both more than twice the rate in London (30.9 per 1000 infant-years, 95% CI 30.4 to 31.3). Nationally, bronchiolitis epidemics peaked once a year, in December (see Figure 4.2). The average weekly variation in observed bronchiolitis admission rates by region are shown in Figure 4.3.

**Birth cohort (2011 to 2016)**


N = 3,808,247

Excluded:

Non-resident or missing information, n = 76,706 (2.0%)

Follow-up time <1 day, n = 4,528 (0.1%)

Children, N = 3,727,013

**Admissions (2012 to 2016)**

Hospital admissions including a diagnosis of bronchiolitis in children <1 year, England, 2012-2016

N = 193,793

Admission within 14 days of discharge from another bronchiolitis admission, n = 16,720 (8.6%)

Admissions, N = 177,073

Removed:

HESID of admission not in birth cohort, n = 21,245 (12.0%)

Duplicate admission, n = 342 (0.2%)

Final cohort for analysis

Children N = 3,727,013

Admissions N = 155,485

**Figure 4.1.** Flow diagram showing derivation of final study cohort and bronchiolitis hospital admissions (linked via unique HESID)
Table 4.1. Cohort characteristics and the number and rate (per 1000 infant-years) of bronchiolitis hospital admissions before 1 year of age among cohort members

<table>
<thead>
<tr>
<th>Region</th>
<th>All infants N</th>
<th>%</th>
<th>Admitted infants* N</th>
<th>Rate (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total</td>
<td>3,727,013</td>
<td>100.0</td>
<td>155,485</td>
<td>50.1 (49.9, 50.4)</td>
</tr>
<tr>
<td>Region</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>North East</td>
<td>176,125</td>
<td>4.7</td>
<td>9,345</td>
<td>63.8 (62.5, 65.1)</td>
</tr>
<tr>
<td>North West</td>
<td>492,007</td>
<td>13.2</td>
<td>28,094</td>
<td>68.7 (67.9, 69.5)</td>
</tr>
<tr>
<td>Yorkshire &amp; the Humber</td>
<td>353,972</td>
<td>9.5</td>
<td>16,838</td>
<td>57.3 (56.4, 58.1)</td>
</tr>
<tr>
<td>East Midlands</td>
<td>289,709</td>
<td>7.8</td>
<td>11,989</td>
<td>49.8 (48.9, 50.7)</td>
</tr>
<tr>
<td>West Midlands</td>
<td>402,365</td>
<td>10.8</td>
<td>19,341</td>
<td>57.9 (57.1, 58.8)</td>
</tr>
<tr>
<td>East of England</td>
<td>403,141</td>
<td>10.8</td>
<td>14,918</td>
<td>44.4 (43.7, 45.1)</td>
</tr>
<tr>
<td>London</td>
<td>705,266</td>
<td>18.9</td>
<td>18,153</td>
<td>30.9 (30.4, 31.3)</td>
</tr>
<tr>
<td>South East</td>
<td>575,762</td>
<td>15.5</td>
<td>22,503</td>
<td>46.9 (46.3, 47.5)</td>
</tr>
<tr>
<td>South West</td>
<td>328,666</td>
<td>8.8</td>
<td>14,304</td>
<td>52.0 (51.2, 52.9)</td>
</tr>
<tr>
<td>Year of birth</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Year</td>
<td>Year of admission</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2011</td>
<td>635,325</td>
<td>17.1</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2012</td>
<td>638,157</td>
<td>17.1</td>
<td>29,853</td>
<td>46.9 (46.4, 47.4)</td>
</tr>
<tr>
<td>2013</td>
<td>621,295</td>
<td>16.7</td>
<td>27,592</td>
<td>44.0 (43.5, 44.6)</td>
</tr>
<tr>
<td>2014</td>
<td>611,217</td>
<td>16.4</td>
<td>28,726</td>
<td>46.6 (46.1, 47.2)</td>
</tr>
<tr>
<td>2015</td>
<td>613,062</td>
<td>16.5</td>
<td>33,286</td>
<td>54.9 (54.3, 55.5)</td>
</tr>
<tr>
<td>2016</td>
<td>607,957</td>
<td>16.3</td>
<td>36,028</td>
<td>58.4 (57.8, 59.1)</td>
</tr>
</tbody>
</table>

*per 1000 infant-years

Figure 4.2. Observed weekly rates (per 1000 infant-years) of hospital admissions for bronchiolitis in England, from January 2012 to December 2016
Figure 4.3. Observed weekly rates of hospital admissions for bronchiolitis, by Region: England, 2012-2016 averaged
4.3.2. Objective 4A: Seasonality of bronchiolitis admissions

Regional level

The estimates of region-specific seasonal parameters derived from the harmonic Poisson regression model assessing bronchiolitis seasonality at the regional level are presented in Table 4.2 and predicted weekly admission rates in Figure 4.4. There was some seasonal variation across regions, with the epidemic peak week ranging from week 50.3 in London (95% CI 50.2 to 50.5) to week 51.4 in the North East (95% CI 51.2 to 51.5). The estimated amplitude at peak epidemic week compared with the mean week was lowest in London ($\hat{\gamma}_j=3.41$, 95% CI 3.31 to 3.52) and highest in the North East ($\hat{\gamma}_j=4.95$, 95% CI 4.72 to 5.19). Each region had an epidemiological duration of 26 or 27 weeks. At week 51, rates of admission were 2.45 and 2.44 times greater in the North West (95% CI 2.39 to 2.52) and the North East (95% CI 2.36 to 2.52) compared with London.

Table 4.2. Derived average annual seasonal estimates following harmonic Poisson regression (Table A4.1, Appendix 4.3), by region

<table>
<thead>
<tr>
<th>Region</th>
<th>Amplitude $\hat{\gamma}^{**}$ (95% CI)</th>
<th>Peak week $\hat{\beta}$ (95% CI)</th>
<th>Phase shift $\hat{\psi}$ (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>North East</td>
<td>4.95 (4.72, 5.19)</td>
<td>51.4 (51.2, 51.5)</td>
<td>-0.21 (-0.23, -0.19)</td>
</tr>
<tr>
<td>North West</td>
<td>4.25 (4.14, 4.36)</td>
<td>50.4 (50.3, 50.5)</td>
<td>-0.34 (-0.35, -0.32)</td>
</tr>
<tr>
<td>Yorkshire and the Humber</td>
<td>4.31 (4.17, 4.46)</td>
<td>50.7 (50.6, 50.8)</td>
<td>-0.29 (-0.31, -0.28)</td>
</tr>
<tr>
<td>East Midlands</td>
<td>4.61 (4.42, 4.80)</td>
<td>51.1 (50.9, 51.9)</td>
<td>-0.25 (-0.27, -0.23)</td>
</tr>
<tr>
<td>West Midlands</td>
<td>4.28 (4.15, 4.42)</td>
<td>50.6 (50.4, 50.7)</td>
<td>-0.31 (-0.32, -0.30)</td>
</tr>
<tr>
<td>East of England</td>
<td>4.89 (4.70, 5.08)</td>
<td>50.8 (50.7, 51.0)</td>
<td>-0.28 (-0.29, -0.26)</td>
</tr>
<tr>
<td>London</td>
<td>3.41 (3.31, 3.52)</td>
<td>50.3 (50.2, 50.5)</td>
<td>-0.34 (-0.36, -0.32)</td>
</tr>
<tr>
<td>South East</td>
<td>4.54 (4.40, 4.68)</td>
<td>50.6 (50.4, 50.7)</td>
<td>-0.31 (-0.32, -0.30)</td>
</tr>
<tr>
<td>South West</td>
<td>4.58 (4.41, 4.75)</td>
<td>51.2 (51.1, 51.4)</td>
<td>-0.23 (-0.25, -0.22)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Region</th>
<th>Duration (weeks)</th>
<th>IRR at week 51 (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>North East</td>
<td>27 (26 to 01)</td>
<td>2.44 (2.36, 2.52)</td>
</tr>
<tr>
<td>North West</td>
<td>26 (25 to 51)</td>
<td>2.45 (2.39, 2.52)</td>
</tr>
<tr>
<td>Yorkshire and the Humber</td>
<td>26 (26 to 52)</td>
<td>2.06 (2.00, 2.12)</td>
</tr>
<tr>
<td>East Midlands</td>
<td>27 (26 to 01)</td>
<td>1.85 (1.79, 1.91)</td>
</tr>
<tr>
<td>West Midlands</td>
<td>27 (25 to 52)</td>
<td>2.08 (2.02, 2.14)</td>
</tr>
<tr>
<td>East of England</td>
<td>26 (26 to 52)</td>
<td>1.68 (1.63, 1.73)</td>
</tr>
<tr>
<td>London</td>
<td>26 (25 to 51)</td>
<td>Reference.</td>
</tr>
<tr>
<td>South East</td>
<td>27 (25 to 52)</td>
<td>1.73 (1.68, 1.73)</td>
</tr>
<tr>
<td>South West</td>
<td>27 (26 to 01)</td>
<td>1.93 (1.87, 1.99)</td>
</tr>
</tbody>
</table>

*Adjusted for year of admission, **amplitude exponentiated
Compared with the average rate of hospital admissions for bronchiolitis in England, predicted rates (derived from the multilevel model, Table A4.2, Appendix 4.3) were highest in CCGs based in the North East and North West and lowest in London and South-East CCGs (Figure 4.5). There was a ratio of 5.3 between the lowest (West Kent) and highest (Stoke on Trent) CCG-based incidence rates. Peak epidemic timing by CCG ranged by 2.9 weeks, from week 49.3 to week 52.2. As illustrated in Figure 4.6, the earliest peaks were seen in North West and South London CCGs and CCGs surrounding London.
Figure 4.5. Predicted peak timing (weeks) of bronchiolitis admission rates, by CCG: England (with London inset)

Figure 4.6. Predicted IRRs of bronchiolitis hospital admissions compared with the national average, by CCG: England (with London inset)
4.3.3. Objective 4B: Multivariable regression models

Results of the multivariable regression models for CCG-level rates of admission and peak timings are displayed in Table 4.3. In mutually adjusted models, a greater IMD score was associated with higher rates of admissions at the CCG level, whilst higher log population density was associated with a slightly lower rate of admissions (Figure 4.7). In total, 23% of the variation in incidence rates at the CCG level was explained by this model, $\eta^2(2)=0.23$ (95% CI 0.13 to 0.31). As shown in Figure 4.8, IMD had a quadratic relationship with peak timing, (likelihood ratio test for the quadratic term, $\chi^2(1)=19.6$, $p<0.001$); both low and high IMD scores were associated with earlier peak timing of bronchiolitis admissions. Higher log population density was associated with earlier peak timing. In total, 38.0% of the variation in peak timing at the CCG level was explained by this model, $\eta^2(2)=0.38$ (95% CI 0.27 to 0.46).

Excluding London-based CCGs

With London-based CCGs excluded from the multivariable regression models (Appendix 4.3, Table A4.3), the adjusted association between IMD and both outcomes (peak timing and IRR of admissions compared with the national average) broadly stayed the same (Table 4.3). The association between log population density and rates of admissions weakened considerably, with only a slight negative association now shown (and only 1% of the variation explained by this variable, $\eta^2(2)=0.01$, 95% CI 0.00 to 0.05). The association between log population density and peak timing decreased from -0.25 (95% CI -0.30 to -0.19) to -0.28 weeks (-0.35 to -0.21).

Table 4.3. Multivariable linear regression analyses assessing the effect of population density and IMD score on IRR compared to the national average and peak timing of CCG epidemic

<table>
<thead>
<tr>
<th></th>
<th>All CCGs*</th>
<th>Excluding London-based CCGs**</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Coefficient (95% CI)</td>
<td>Partial $\eta^2$ (95% CI)</td>
</tr>
<tr>
<td>IRR</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Population density†</td>
<td>-0.09 (-0.12, -0.05)</td>
<td>0.11 (0.04, 0.20)</td>
</tr>
<tr>
<td>IMD</td>
<td>0.02 (0.02, 0.03)</td>
<td>0.22 (0.13, 0.31)</td>
</tr>
<tr>
<td>Constant</td>
<td>1.19 (0.98, 1.40)</td>
<td>0.75 (0.53, 0.97)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Peak timing (weeks)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Population density†</td>
<td>-0.25 (-0.30, -0.19)</td>
<td>0.30 (0.20, 0.39)</td>
</tr>
<tr>
<td>IMD</td>
<td>0.11 (0.07, 0.14)</td>
<td>0.17 (0.09, 0.26)</td>
</tr>
<tr>
<td>IMD²</td>
<td>-0.002 (-0.002, -0.001)</td>
<td>0.10 (0.04, 0.18)</td>
</tr>
<tr>
<td>Constant</td>
<td>50.87 (50.40, 51.34)</td>
<td>51.08 (50.49, 51.67)</td>
</tr>
</tbody>
</table>
*IRR and peak week derived from harmonic multilevel mixed-effect Poisson regression analysis displayed in Table A4.2 (Appendix 4.3); ** IRR and peak week derived from harmonic multilevel mixed-effect Poisson regression analysis displayed in Table A4.3 (Appendix 4.3); †Log-scale

**Figure 4.7.** Observed and predicted variation in CCG-level rates of bronchiolitis hospital admissions compared with the national average, by IMD score and log population density (including London-based CCGs)

**Figure 4.8.** Observed and predicted peak timing of CCG-level bronchiolitis hospital admissions, by IMD score and log population density (including London-based CCGs)
4.3.4. Sensitivity analysis

To assess whether the uncertainty of the stage 1 estimation had an impact on inferences at stage 2, I calculated bootstrapped standard errors on 20% of the total sample and compared them to those obtained without bootstrap on the same data. The results showed very minimal differences, supporting the interpretations above, especially given the size of the whole sample (see Table A4.4, Appendix 4.4). The Moran’s I values were positive for the IRR residuals ($I=0.28$) and the peak timing residuals ($I=0.37$) of spatially proximal CCGs. When I included only the first admission by each child in the analyses, regional amplitudes increased and peak epidemic timing was approximately 0.2 weeks earlier (Table A4.5, Appendix 4.4). Relative differences in the seasonality of admissions across regions and CCG-level analyses remained broadly the same (Tables A4.6 and A4.7, Appendix 4.4).

4.4. Discussion

I found large variation in the size of the epidemic peak and accompanying admission rates for bronchiolitis, particularly when comparing London to the rest of the country. At week 51 (mid-December) rates of admissions were 2.4 to 2.5 times higher in Northern regions of the country compared with London. There was a 5.3-fold difference in rates across CCGs, with about a quarter of the difference explained by area-level deprivation. The shape of the epidemic curve was similar across regions, with admission rates increasing continually for 26 or 27 weeks of the year and epidemics peaking between week 50 and 51 (2nd and 3rd weeks of December). At the CCG-level the difference in peak week increased to 3 weeks, with earlier peaks tending to occur in areas with higher population densities, such as London and Manchester.

4.4.1. Strengths and weaknesses

The birth cohort was created from HES APC, a national administrative dataset of hospital admissions in England. Data in HES APC are recorded by clinical coders in each hospital and coding consistency may, therefore vary by hospital according to diagnostic protocols and recording practices. Diagnostic codes are rarely entered in accident and emergency departments in England, and there is no national primary care database covering all English general practices. Therefore, I could not examine bronchiolitis seasonality in accident and emergency attendances or general practice consultations, which, it could be hypothesised, may peak before hospital admissions. Private hospital activity amongst non-NHS funded patients is not captured in HES APC. However, since emergency treatment is not typically covered under private medical insurance in the UK and NHS care is (for the most part) free at the point of need, I expect that very few children are admitted privately for acute illnesses like
bronchiolitis. Overall, with a large number of bronchiolitis admissions across the whole of England, this study provides a contextual understanding of the seasonality of this infection and how it varies across the country.

I modelled admissions using harmonic functions, which allowed me to assess variability in the seasonality of bronchiolitis by area of the country. The predicted regional model produced seasonal patterns similar to the observed rates of admissions but underestimated the amplitude of the regional epidemics. Fitting splines may have improved estimation but would not have allowed me to straightforwardly estimate seasonal parameters. Using a single pair of sine/cosine terms in the model mirrored the single annual peak exhibited in the data, and enabled me to estimate the peak epidemic week seen each year in England. I was unable to account for uncertainty in the estimates of incidence rates and peak timing in the second part of the CCG-level model; however, the results were shown to be robust in sensitivity analyses.

4.4.2. Research findings: comparison with other studies

Objective 4A: Seasonality of bronchiolitis admissions across England

As expected, a substantial single epidemic peak in hospital admissions for bronchiolitis occurred each winter in England between 2012 and 2016 in mid-December. A small increase in hospital admissions around the fourth week of year can also be seen in Figures 4.2 and 4.3. It is likely that this increase in cases is driven by the post-holiday return to day care and/or school (by older siblings), which are known sources of infant RSV infection. The 5.3-fold geographical variation in bronchiolitis admission rates by CCG found here matches Green et al.’s analysis by Local Government Area between 1999 and 2011. Another analysis that looked at Primary Care Trusts estimated that the variation was almost 3 times higher than in my study, although this difference can be explained by their definition of bronchiolitis. Children up to the age of 2 admitted to hospital with a primary diagnoses of bronchiolitis were included in their definition, leading to some area-level rates lower than 5 per 1000 children. I add to these previous findings with information about the epidemic curves, which were found to have a similar duration across regions of England but differ with regards to the peak week at the CCG-level.

Objective 4B: Association between CCG-level epidemic timing, rates of admissions and demographic characteristics

Similar to findings from previous research, I found that one fifth of the variation in admission rates across England was explained by socioeconomic factors, after controlling for population density. Socioeconomic deprivation is a broad facet associated with risk factors for severe
RSV infection including tobacco smoke exposure, housing conditions and overcrowding, amongst others.\textsuperscript{4,45,139} I used a small-area level indicator of deprivation, IMD, to capture socio-economic deprivation of the children contributing to this study. IMD is widely used in studies of health outcomes in England, and has been shown to be associated with bronchiolitis admission rates.\textsuperscript{4,344} Further research is required to establish which particular aspects of deprivation have the highest impact on bronchiolitis admissions.

Converse to expectation, larger population density was associated with lower admission rates; however, sensitivity analyses showed that this relationship was driven by London based CCGs. The difference in emergency admission rates in London compared with the rest of England is stark, but not a new finding—having previously been reported for a range of conditions.\textsuperscript{45,344,345} Plausible reasons for differential admission rates include primary care accessibility and availability, and differences in admission thresholds.\textsuperscript{4} Data published by Public Health England show that, whilst rates of emergency admissions for children aged 0-4 are low in London, accident and emergency (A&E) rates are higher than the national average, potentially reflecting regional differences in admission practices.\textsuperscript{347} This hypothesis is corroborated by early results from research using English A&E linked to HES APC data led by my colleague, Selina Nath.\textsuperscript{348}

My results suggest that population density is a stronger predictor of epidemic peak timing than of incidence rates, even after adjustment for IMD. Similar to the results of an RSV epidemic study in Connecticut,\textsuperscript{124} I found that higher population density (residents per km\textsuperscript{2}) was associated with an earlier peak timing. Area-level socioeconomic deprivation had an inverted U-shaped association with peak timing, which may be indicative of shared factors between CCGs on the extremes of socioeconomic deprivation. Figure 4.6 illustrates that the earliest peaks are in South London, Manchester and Birmingham—all urban areas with major national and international transport links. High levels of travel, and increased contact and mixing amongst the population likely play a role in viral spread in these areas. Pitzer at al.\textsuperscript{111} undertook detailed modelling of RSV epidemics across USA, noting associations between vapour pressure, temperature, precipitation and the timing of epidemics. Notably, even in a country with much greater climatic variation that the UK, Pitzer at al.\textsuperscript{111} were unable to account for the finding that RSV activity begins in Florida, leading the authors to contemplate the potential role of population mixing.

4.4.3. Implications of study findings

The findings have implications on preparedness for bronchiolitis infections. Given the gap in timing between peak epidemics across the country, messaging systems could be enacted to
relay warnings from CCGs experiencing early peaked admissions to other areas. These systems could inform the distribution of prophylaxis or future vaccinations, and be used to time simple public health messages, such as warning families to avoid taking their young infants into public spaces or encourage frequent handwashing for older siblings. The results also show that surge capacity plans for paediatric intensive care units would likely profit from real-time information on hospital admissions for bronchiolitis across the country. This is particularly important given that RSV infections are not currently captured in real-time in the UK.

Green et al. found that admissions for bronchiolitis have risen substantially over the last few decades and my updated analyses show a continuation of this trend. As described elsewhere, it is likely that changes in admission thresholds and accessibility of primary care services have driven this increase rather than disease severity or increased incidence of bronchiolitis. By 2016, the admission rate among infants in this study had increased to 58.4 per 1000 infant-years (95% CI 57.8 to 59.1), highlighting the need to focus resources on research and interventions aimed at this condition. Evaluations of alternative models of care that reduce the need for hospital admission for acutely ill infants is of particular importance. Future geospatial studies would profit from the inclusion of factors unavailable in the dataset, such as a measure of admission thresholds and population mixing, to examine these factors further. Additional investigation of the incongruent patterns of admissions in London compared with the rest of the country is particularly warranted, as well as examining the impact of ecological factors such as climate and air pollution on the timing of admission rates.

4.5. Chapter summary

This nationwide descriptive analysis of bronchiolitis admissions has shown variation in the size of the epidemic peak across England after accounting for seasonal components of admission rates. At the regional level, predicted rates of admissions were almost 2.5 times greater in the North of England compared to London. The duration of the epidemic was similar across the country, although there was a 3-week difference in the peak week of admissions at the CCG-level. Approximately one quarter of the variation in admission rates and two-fifths of the variation in peak timing of hospital admissions for bronchiolitis were explained by local demographic and socio-economic characteristics. Implementation of an early warning system could help to prepare hospitals for peak activity and to time public health messages.
Chapter 5. Is socioeconomic position associated with bronchiolitis seasonality?

Recap and chapter overview

Chapter 4 investigated how the timing and volume of hospital admissions for bronchiolitis is associated with demographic characteristics, including area-level socioeconomic deprivation, across England. In this Chapter, I use the same English longitudinal dataset to examine the interaction between seasonality in weekly hospital admissions for bronchiolitis and socioeconomic position at the individual level. This work is the second study to address thesis objective 1:

*To describe the association between seasonality, early life socioeconomic position and bronchiolitis admission rates among infants across England.*

A research paper based on the content of this chapter has been published in the Journal of Epidemiology and Community Health (included in Appendix 5.1).

5.1. Background

As shown in Chapter 4, there is a sharp peak in hospital admissions for bronchiolitis in England in December, indicating an increased burden on health care services during the winter months.\(^4,16,351\) The main pathogen causing bronchiolitis is respiratory syncytial virus (RSV); a virus that has a diverse pattern of seasonal changes globally, but is far more prevalent during winter months in temperate Northern climates such as the UK.\(^6,109,110\) Although the precise mechanisms driving this seasonal pattern are yet to be determined, several interacting virus and host (i.e. human) factors have been proposed.\(^2,352\) In temperate climates, RSV tend to survive longer in the lower temperatures observed in wintertime, with higher relative humidity also thought to play a role.\(^352\) In parallel, this is a period where humans spend more time indoors together, thereby increasing the likelihood of virus transmission. It is also thought that humans are more susceptible to infection at this time owing to a combination of factors including respiratory tract dehydration and vitamin D deficiency.\(^77,353\)

Research conducted in England and Scotland has shown that children from lower socioeconomic groups are at increased risk of admission to hospital for bronchiolitis.\(^4,84,354\) However, it is not known whether there are differences in the seasonal pattern of bronchiolitis admissions across socioeconomic groups. It is plausible that, due to social patterning of
factors that affect RSV transmission and infection—including damp, overcrowded housing conditions and a greater prevalence of underlying health problems—there may also be differential seasonal patterns in hospital admissions for bronchiolitis. Understanding these patterns could help to inform preventive interventions to reduce bronchiolitis admissions, including optimal timing of palivizumab prophylaxis prescribing in different population groups, or targeting of maternal RSV vaccination, when this becomes available in the future.

The aim of this Chapter is to determine how socioeconomic position is associated with the seasonality of hospital admissions for bronchiolitis amongst infants in England.

There are two objectives:

**Objective 5A** To calculate the proportion of all infant hospital admissions in England between 2012 and 2016 attributable to bronchiolitis by year, month and socioeconomic group.

**Objective 5B** To quantify the extent to which seasonality of bronchiolitis admissions is modified by socioeconomic position.

### 5.2. Methods

#### 5.2.1. Data sources and study cohort

The data source for this study was hospital episode statistics admitted patient care (HES APC) linked to Office for National Statistics (ONS) mortality records as presented in Chapters 3 and 4. I used the same birth cohort as in Chapter 4 (see Section 4.2.1). Briefly, this consisted of children born from 1st January 2011 to 31st December 2016, who were followed from birth or the 1st of January 2012, whichever occurred last. Follow up was until their first birthday, date of migration, 31st December 2016 or date of death, whichever occurred first. Using the infant’s unique HESID, data on admissions to hospital within the first year of life were added to birth data. Infants were excluded from cohort analyses if they had missing information, were from a multiple birth or were stillborn. Infants with a non-English address recorded at birth (identified through lower super output area, LSOA) were also excluded to prevent potential loss to follow up. Where a non-English LSOA was recorded in a subsequent infant admission, a censoring date was placed halfway between the date of the infant’s last known resident admission and the non-resident admission.

#### 5.2.2. Study outcome

I identified hospital admissions with a diagnosis of bronchiolitis in HES APC using the International Classification of Diseases version 10 (ICD-10) code J21 for acute bronchiolitis.
As explained in Chapter 4 (Section 4.2.2), all J21 subcategories were included in this definition because of the low sensitivity of RSV-specific ICD-10 codes. All admissions with bronchiolitis recorded as either the primary or secondary diagnosis for an infant during their first year of life were included in the analyses. To calculate admission rates, any admission for bronchiolitis within 14 days of discharge from another bronchiolitis admission was assumed to be associated with the same infection, and therefore only the first of these admissions was included in the analyses.

The numerator for calculating the total burden of infant admissions attributable to bronchiolitis included all admissions with bronchiolitis recorded as either the primary diagnosis or one of the secondary diagnoses where the patient was ≤365 days old. Any admission ≥1 day after discharge from a previous admission was counted as a separate admission. To ascertain a denominator for the calculation of the total burden of infant admissions attributable to bronchiolitis, I extracted all emergency and planned hospital admissions (excluding birth admissions) between 1st January 2012 and 31st December 2016 where the patient was ≤365 days old.

### 5.2.3. Covariates

I used LSOA-level IMD 2015 to capture socioeconomic position for each cohort member. IMD was derived from mother’s recorded LSOA at the time of delivery or the earliest mention of LSOA in infant’s hospital admission. IMD quintiles were used for analyses, which the Ministry of Housing, Communities and Local Government (MHCLG) calculates by ranking LSOAs in England in order of IMD score and dividing them into 5 equal groups. In my analyses, the lowest ranked quintile represents the least deprived and the highest ranked quintile the most deprived fifth of LSOAs in England.

Health and disability is one of the seven domains used to calculate IMD 2015 scores. This domain is constructed from four indicators, two of which (acute morbidity and mood and anxiety disorders) include hospital admission statistics. Given that my outcome is also based on hospital admission data, this could be a source of bias in my study. I therefore also created IMD 2015-minus the health domain-quintiles to look at the difference in results when the health domain is excluded from IMD. The MHCLG have a purposefully created dataset that enables researchers to recalculate IMD with the exclusion of specific domains. The scores in this dataset have been standardised by ranking, and then transformed to an exponential distribution. Following the guidance issued by the MHCLG, I combined the standardised domain scores with all domain weights (shown in Table 5.1) apart from health for each LSOA as follows:
(income deprivation domain score*domain-weight) + (employment deprivation domain score*domain-weight) + (education, skills and training deprivation domain score*domain-weight) + (barriers to housing and services deprivation domain score*domain-weight) + (crime deprivation domain score*domain-weight) + (living environment deprivation domain score*domain-weight)

I then ranked the scores from lowest to highest and divided them into 5 equal groups to create quintiles. The correlation coefficient between IMD scores with and without the health domain was 0.995.

**Table 5.1. IMD 2015 domain weightings**

<table>
<thead>
<tr>
<th>Deprivation domain</th>
<th>IMD domain weighting (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Income</td>
<td>22.5</td>
</tr>
<tr>
<td>Employment</td>
<td>22.5</td>
</tr>
<tr>
<td>Health and disability</td>
<td>13.5</td>
</tr>
<tr>
<td>Education, skills and training</td>
<td>13.5</td>
</tr>
<tr>
<td>Barriers to housing and service</td>
<td>9.3</td>
</tr>
<tr>
<td>Crime</td>
<td>9.3</td>
</tr>
<tr>
<td>Living environment</td>
<td>9.3</td>
</tr>
</tbody>
</table>

Infant sex (male or female), month and year of birth were extracted from infant’s birth record. Infant’s age at admission for bronchiolitis was calculated by subtracting the bronchiolitis admission date from the infant’s admission date in their birth episode. Age was split into three groups: <3 months, 3 to <6 months and 6 to <12 months. Government Office Region of residence was used to indicate the region of England within which the infant lived at time of birth. HES APC includes information on additional risk factors such as gestational age, mode of delivery and congenital anomalies; however, as these variables are likely on the causal pathway between socioeconomic position and admission for bronchiolitis, they were not included in this analyses. Inclusion of mediators in associational analyses would introduce overadjustment bias into the model.

5.2.4. Statistical methods

Stata 15.0 was used for data analysis and Microsoft Excel 2013 to create graphs. The Stata code for the harmonic regression analysis can be found in Appendix 5.2.

Bronchiolitis admission rates across the cohort were calculated by dividing the number of new admissions for bronchiolitis by person-time at risk for all infants at risk in the birth cohort, and are expressed as annual admission-based rates per 1000 infant-years. I also calculated person-based rates where, for infants with multiple admissions, only the first admission was counted.
**Objective 5A: the proportion of infant hospital admissions attributable to bronchiolitis**

To calculate the proportion of all infant hospital admissions attributed to bronchiolitis between 2012 and 2016, I divided the number of admissions with a diagnosis of bronchiolitis by the total number of hospital admissions. Admission data was split by year and month of admission and IMD quintile at time of admission. Birth admissions and infants with a non-English LSOA at the time of admission to hospital were excluded from the analysis.

**Objective 5B: the extent to which seasonality of bronchiolitis admissions is modified by socioeconomic position**

Event rates were modelled using a Poisson regression model with robust standard errors to account for multiple admissions for the same child. As in Chapter 4, I did not use negative binomial models because there was no evidence of over-dispersion in events. The impact of seasonality on rates of admission was modelled using a harmonic function of time in weeks using the same method described in section 4.2.4. I also included interaction terms of IMD quintile with the sine and cosine regression coefficients (denoted \( \delta_j \) and \( \delta_j \)) to assess evidence of effect modification of IMD quintile by seasonality, formally tested using a Wald \( \chi^2 \) test. Year of admission, region, sex, month of birth, age group and interaction terms of age group with the sine and cosine regression coefficients were selected a priori as covariates in the model to increase precision given their known associations with the outcome as identified in the literature. See Box 5.1 for the full model specification.

Using the estimated model coefficients, I calculated the amplitude (log) (\( \gamma_j \)), phase (\( \psi_j \)) and peak week of the epidemic curve of each IMD quintile at reference values of the other covariates using the formulae outlined in Section 4.2.4. I also calculated the incidence rate ratios (IRR) at the average peak week of the top four IMD quintiles relative to the lowest IMD quintile. The delta method, as implemented by the Stata command `lincom`, was used to calculate 95% confidence intervals for each parameter. Epidemic duration for each IMD quintile was calculated using the same method outlined in Chapter 4 (from the first of three consecutive weeks with increasing predicted rates to the first of three consecutive weeks with decreasing predicted rates).

**Sensitivity analysis**

To assess whether inclusion of hospital admission statistics in the measure of socioeconomic position in this study (IMD) influences the results, I replicated the harmonic regression analysis with IMD 2015-minus the health domain-quintiles.
Box 5.1. Model specifications

Let \( n_{jk}(t) \), the number of bronchiolitis admissions in IMD group \( j \) and year \( k \) observed at time \( t \) (measured in weeks), follow a Poisson distribution with rate \( \lambda_{jk}(t) = E(n_{jk}(t))/N_{jk}(t) \) where \( N_{jk}(t) \) denote the person-time at risk in IMD group \( j \) and year \( k \) at time \( t \). I modelled this rate, after log-transformation, as a function of year and time of admission as follows:

\[
\log(\lambda_{jk}(t)) = \beta_0 + \beta_1 \sin(2\pi t/T) + \beta_2 \cos(2\pi t/T) + \sum_{j=1}^{5} \alpha_j I_{\text{IMD}=j} \\
+ \sum_{j=1}^{5} \delta_{1j} \sin(2\pi t/T) I_{\text{IMD}=j} + \sum_{j=1}^{5} \delta_{2j} \cos(2\pi t/T) I_{\text{IMD}=j} + \sum_{k=1}^{3} \theta_k I_{\text{age}=k} \\
+ \sum_{k=1}^{3} \theta_k \sin(2\pi t/T) I_{\text{age}=k} + \sum_{k=1}^{3} \theta_k \cos(2\pi t/T) I_{\text{age}=k} + \sum_{k=1}^{K} \theta_k I_{X=x}
\]

Where: \( T \) is the length of period within one harmonic cycle (i.e. 1 year = 52.14 weeks); and \( I_{X=x} \) is the binary indicator of the variable \( X \) taking value \( x \); \( K \) indicates covariates included in the model (e.g. age, sex and year categories). The parameter \( \beta_0 \) is the intercept, \( \beta_1 \) and \( \beta_2 \) are harmonic function coefficients, \( \delta_{1j} \) and \( \delta_{2j} \) are IMD group-specific harmonic function coefficients, and the parameters \( \alpha_1, \delta_{11}, \delta_{21} \) and \( \theta_1 \) are all constrained to be zero to deal with the collinearity of the binary indicators.
5.3. **Results**

5.3.1. **Objective 5A: the proportion of admissions attributable to bronchiolitis**

The average annual proportion of all infant hospital admissions in England with a diagnosis of bronchiolitis was 15.0% over the five study years (Table 5.2). The proportion increased over time, from 14.2% in 2012 to 16.6% in 2016. The same pattern, at a slightly lower percentage, was present when only hospital admissions with a primary diagnosis of bronchiolitis were included (12.9% in 2012, rising to 14.9% in 2016). The burden of bronchiolitis was concentrated in the winter months. In December 2012 to 2016 combined, 39.8% of total infant admissions had a diagnosis of bronchiolitis recorded compared to 2.3% of August admissions over the same period. Including only primary diagnoses reduced this proportion slightly; 36.4% of all infant admissions in December and 2.0% in August had primary diagnoses of bronchiolitis.

Raw numbers of admissions, overall and with a diagnosis of bronchiolitis, were far higher in the more deprived compared to the less deprived IMD quintiles. Of admissions with a recorded IMD score, 30.5% overall and 33.1% admissions with a diagnosis of bronchiolitis were from infants in the most deprived fifth of LSOAs in England. Proportionally, a slight graded pattern in admissions with a bronchiolitis diagnosis was shown between IMD quintiles, from 14.0% in the least deprived to 16.3% in the most deprived group for admissions with any diagnosis of bronchiolitis.

**Table 5.2.** Postnatal hospital admissions in children <1 year old, any condition and bronchiolitis-related, by year, month and IMD quintile: England, 2012-2016

<table>
<thead>
<tr>
<th></th>
<th>Admissions for any condition</th>
<th>Bronchiolitis admissions</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>N (% of total)</td>
<td>Primary or secondary diagnosis</td>
</tr>
<tr>
<td>Overall (2012-2016)</td>
<td>1,292,649 (100.0)</td>
<td>193,793 (15.0)</td>
</tr>
<tr>
<td>Year</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2012</td>
<td>262,411 (20.3)</td>
<td>37,136 (14.2)</td>
</tr>
<tr>
<td>2013</td>
<td>252,053 (19.5)</td>
<td>34,264 (13.6)</td>
</tr>
<tr>
<td>2014</td>
<td>250,454 (19.4)</td>
<td>35,982 (14.4)</td>
</tr>
<tr>
<td>2015</td>
<td>255,866 (19.8)</td>
<td>41,408 (16.2)</td>
</tr>
<tr>
<td>2016</td>
<td>271,865 (21.0)</td>
<td>45,003 (16.6)</td>
</tr>
<tr>
<td>Month</td>
<td></td>
<td></td>
</tr>
<tr>
<td>January</td>
<td>112,008 (8.7)</td>
<td>24,116 (21.5)</td>
</tr>
<tr>
<td>February</td>
<td>104,485 (8.1)</td>
<td>13,787 (13.2)</td>
</tr>
<tr>
<td>March</td>
<td>113,802 (8.8)</td>
<td>11,647 (10.2)</td>
</tr>
<tr>
<td>April</td>
<td>101,518 (7.9)</td>
<td>8,077 (8.0)</td>
</tr>
<tr>
<td>May</td>
<td>100,649 (7.8)</td>
<td>6,238 (6.3)</td>
</tr>
<tr>
<td>June</td>
<td>93,729 (7.3)</td>
<td>4,004 (4.3)</td>
</tr>
</tbody>
</table>
Chapter 5 - Bronchiolitis seasonality: interaction with SEP

<table>
<thead>
<tr>
<th>Month</th>
<th>Cases</th>
<th>Rate (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>July</td>
<td>96,421</td>
<td>3,432 (3.6)</td>
</tr>
<tr>
<td>August</td>
<td>85,770</td>
<td>1,707 (2.0)</td>
</tr>
<tr>
<td>September</td>
<td>90,995</td>
<td>4,767 (5.2)</td>
</tr>
<tr>
<td>October</td>
<td>109,270</td>
<td>12,823 (11.7)</td>
</tr>
<tr>
<td>November</td>
<td>135,434</td>
<td>38,268 (28.3)</td>
</tr>
<tr>
<td>December</td>
<td>148,568</td>
<td>54,122 (36.4)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>IMD quintile</th>
<th>Cases</th>
<th>Rate (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 Least deprived</td>
<td>173,847</td>
<td>22,239 (12.8)</td>
</tr>
<tr>
<td>2</td>
<td>195,163</td>
<td>25,415 (13.0)</td>
</tr>
<tr>
<td>3</td>
<td>229,939</td>
<td>30,164 (13.1)</td>
</tr>
<tr>
<td>4</td>
<td>287,007</td>
<td>39,185 (13.7)</td>
</tr>
<tr>
<td>5 Most deprived</td>
<td>388,925</td>
<td>57,215 (14.7)</td>
</tr>
<tr>
<td>Missing</td>
<td>17,318</td>
<td>1,345 (7.8)</td>
</tr>
</tbody>
</table>

5.3.2. Cohort description

Of the 3,808,247 singleton children born between 2011 and 2016, 81,234 (2.1%) were excluded from the final study cohort as shown in Figure 5.1. This included 20,905 (0.5%) infants excluded due to non-resident status, 55,025 (1.4%) infants with missing information for region, IMD and/or sex and 4,528 (0.1%) infants with follow-up less than 1 day. The final cohort comprised 3,727,789 singleton infants, of which 48.7% were female, 27.2% were in the most deprived quintile of IMD and 18.9% were born to mothers residing in London (Table 5.3). There are 776 more children included in this cohort compared to Chapter 4 due to slight differences in exclusions due to missing data. The number and rate of hospital admissions for bronchiolitis \( N = 155,485 \), 50.1 per 1000 infant-years, 95% CI 49.9 to 50.4) and average follow-up time per child (305 days) remained the same. The overall infant-based admission rate was 46.2 per 1000 infant-years (95% CI 46.0 to 46.5).

Alongside later year of admission and residence in the North West or North East of England as discussed in Chapter 4 (see Table 4.1), rates of admissions for bronchiolitis were highest in the most socioeconomically deprived groups, males, and younger infants. The average admission rate was 78.9 per 1000 infant-years (95% CI 77.9 to 80.0) for infants born in October compared to 33.7 per 1000 infant-years (95% CI 33.0 to 34.4) for infants born in March. Figure 5.2 displays observed weekly rates of admissions by IMD, illustrating differential rates of admissions by level of socioeconomic position across the year and a clear annual peak in December.
**Figure 5.1.** Flow diagram to show study participant and hospital admission selection
Table 5.3. Distribution of births and bronchiolitis admissions (per 1000 infant-years) in the cohort

<table>
<thead>
<tr>
<th>Variable</th>
<th>Infants N</th>
<th>%</th>
<th>Bronchiolitis admissions</th>
<th>Person-based*</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>Admission-based N</td>
<td>Rate (95% CI)</td>
</tr>
<tr>
<td>Total</td>
<td>3,727,789</td>
<td>100.0</td>
<td>155,485</td>
<td>50.1 (49.9, 50.4)</td>
</tr>
<tr>
<td>IMD quintiles</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 Least deprived</td>
<td>551,740</td>
<td>14.8</td>
<td>18,877</td>
<td>41.1 (40.5, 41.7)</td>
</tr>
<tr>
<td>2</td>
<td>615,852</td>
<td>16.5</td>
<td>22,379</td>
<td>43.6 (43.1, 44.2)</td>
</tr>
<tr>
<td>3</td>
<td>700,171</td>
<td>18.8</td>
<td>27,051</td>
<td>46.4 (45.8, 46.9)</td>
</tr>
<tr>
<td>4</td>
<td>846,766</td>
<td>22.7</td>
<td>35,015</td>
<td>49.7 (49.2, 50.2)</td>
</tr>
<tr>
<td>5 Most deprived</td>
<td>1,013,260</td>
<td>27.2</td>
<td>52,163</td>
<td>62.0 (61.5, 62.5)</td>
</tr>
<tr>
<td>Sex</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Female</td>
<td>1,813,651</td>
<td>48.7</td>
<td>61,349</td>
<td>40.8 (40.4, 41.1)</td>
</tr>
<tr>
<td>Male</td>
<td>1,914,138</td>
<td>51.4</td>
<td>94,134</td>
<td>59.3 (59.0, 59.7)</td>
</tr>
<tr>
<td>Month of birth</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>January</td>
<td>310,594</td>
<td>8.3</td>
<td>9,362</td>
<td>36.3 (35.6, 37.1)</td>
</tr>
<tr>
<td>February</td>
<td>284,912</td>
<td>7.6</td>
<td>8,022</td>
<td>33.9 (33.1, 34.6)</td>
</tr>
<tr>
<td>March</td>
<td>305,605</td>
<td>8.2</td>
<td>8,536</td>
<td>33.7 (33.0, 34.4)</td>
</tr>
<tr>
<td>April</td>
<td>298,220</td>
<td>8.0</td>
<td>8,847</td>
<td>35.8 (35.1, 36.5)</td>
</tr>
<tr>
<td>May</td>
<td>316,825</td>
<td>8.5</td>
<td>10,283</td>
<td>39.0 (38.3, 39.8)</td>
</tr>
<tr>
<td>June</td>
<td>310,086</td>
<td>8.3</td>
<td>11,140</td>
<td>43.4 (42.6, 44.2)</td>
</tr>
<tr>
<td>July</td>
<td>325,966</td>
<td>8.7</td>
<td>13,636</td>
<td>50.4 (49.6, 51.3)</td>
</tr>
<tr>
<td>August</td>
<td>319,654</td>
<td>8.6</td>
<td>15,767</td>
<td>59.3 (58.4, 60.2)</td>
</tr>
<tr>
<td>September</td>
<td>322,995</td>
<td>8.7</td>
<td>18,980</td>
<td>70.6 (69.6, 71.6)</td>
</tr>
<tr>
<td>October</td>
<td>322,920</td>
<td>8.7</td>
<td>21,306</td>
<td>78.9 (77.9, 80.0)</td>
</tr>
<tr>
<td>November</td>
<td>304,734</td>
<td>8.2</td>
<td>17,568</td>
<td>69.0 (68.0, 70.1)</td>
</tr>
<tr>
<td>December</td>
<td>305,278</td>
<td>8.2</td>
<td>12,038</td>
<td>46.9 (46.1, 47.8)</td>
</tr>
<tr>
<td>Age at event</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>&lt; 3 months</td>
<td>61,926</td>
<td>80.2</td>
<td>80.2 (79.5, 80.8)</td>
<td>60,500</td>
</tr>
<tr>
<td>3 to &lt; 6 months</td>
<td>45,907</td>
<td>59.3</td>
<td>59.3 (58.7, 59.8)</td>
<td>41,785</td>
</tr>
<tr>
<td>6 to &lt; 12 months</td>
<td>47,652</td>
<td>50.7</td>
<td>50.7 (50.4, 51.0)</td>
<td>40,921</td>
</tr>
<tr>
<td>Month of admission</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>January</td>
<td>19,154</td>
<td>72.7</td>
<td>72.7 (71.7, 73.7)</td>
<td>17,331</td>
</tr>
<tr>
<td>February</td>
<td>10,968</td>
<td>45.4</td>
<td>45.4 (44.6, 46.3)</td>
<td>9,249</td>
</tr>
<tr>
<td>March</td>
<td>9,303</td>
<td>35.3</td>
<td>35.3 (34.6, 36.1)</td>
<td>7,599</td>
</tr>
<tr>
<td>April</td>
<td>6,536</td>
<td>25.6</td>
<td>25.6 (25.0, 26.3)</td>
<td>6,497</td>
</tr>
<tr>
<td>May</td>
<td>5,094</td>
<td>19.3</td>
<td>19.3 (18.8, 19.9)</td>
<td>4,774</td>
</tr>
<tr>
<td>June</td>
<td>3,246</td>
<td>12.8</td>
<td>12.8 (12.3, 13.2)</td>
<td>2,910</td>
</tr>
<tr>
<td>July</td>
<td>2,801</td>
<td>10.6</td>
<td>10.6 (10.3, 11.0)</td>
<td>2,470</td>
</tr>
<tr>
<td>August</td>
<td>1,575</td>
<td>6.0</td>
<td>6.0 (5.7, 6.3)</td>
<td>1,401</td>
</tr>
<tr>
<td>September</td>
<td>4,419</td>
<td>17.4</td>
<td>17.4 (16.9, 17.9)</td>
<td>3,938</td>
</tr>
<tr>
<td>October</td>
<td>11,593</td>
<td>44.1</td>
<td>44.1 (43.3, 44.9)</td>
<td>10,761</td>
</tr>
<tr>
<td>November</td>
<td>33,838</td>
<td>133.8</td>
<td>133.8 (132.4, 135.2)</td>
<td>32,123</td>
</tr>
<tr>
<td>December</td>
<td>46,958</td>
<td>177.7</td>
<td>177.7 (176.1, 179.3)</td>
<td>44,153</td>
</tr>
</tbody>
</table>

*where only the first of multiple hospital admissions for bronchiolitis by the same child is counted.
Objective 5B: effect modification by socioeconomic position

The multivariable model that included interaction terms between IMD and the two harmonic functions fitted the data better than the model without interactions, Wald test, $\chi^2(8) = 239.2$, $p<0.001$ (Table A5.1, Appendix 5.3). Fitted values and seasonal estimates from the model are presented in Figure 5.3 and Table 5.4. The amplitude (that is, the difference in rates of bronchiolitis hospital admissions between the peak and trough of the epidemic curve) was smallest in the most deprived IMD quintiles and largest in the least deprived quintiles. This points to less seasonal variation in the rates of admissions in the more disadvantaged groups. The average peak timing of the annual epidemic varied marginally across IMD quintiles, from week 49.4 (95% CI 49.2 to 49.5) in the least deprived quintile to week 50.3 (95% CI 50.1 to 50.4) in the most deprived quintile. The estimated epidemic duration was 26 weeks across all IMD quintiles. After adjustment for covariates, infants in the most deprived group had a 40% (IRR 1.40, 95% CI 1.37 to 1.43) greater risk of admission to hospital for bronchiolitis at week 50 compared to infants in the least deprived group.
5.3.4. Sensitivity analysis: excluding the health domain from IMD

There was very little difference between the seasonal estimates when the analysis was replicated using IMD-excluding health quintiles. The amplitude was almost identical across quintiles and the phase shift and peak week estimates remained the same. The point estimates of the IRR of admission rates at week 50, compared to the least deprived quintile, became slightly smaller (moving from a 40% to 37% increased risk in the most extreme example) but confidence interval overlapped considerably.

Table 5.4. Average annual seasonal estimates derived from harmonic Poisson regression models (see Appendix 4.3), by IMD quintile

<table>
<thead>
<tr>
<th>IMD quintile</th>
<th>Amplitude* (95% CI)</th>
<th>Phase shift (95% CI)</th>
<th>Peak week (95% CI)</th>
<th>IRR at week 50 (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Full IMD</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1-Least deprived</td>
<td>3.97 (3.81, 4.12)</td>
<td>-0.45 (-0.47, -0.44)</td>
<td>49.4 (49.2, 49.5)</td>
<td>Reference</td>
</tr>
<tr>
<td>2</td>
<td>3.83 (3.70, 3.97)</td>
<td>-0.41 (-0.43, -0.39)</td>
<td>49.7 (49.6, 49.9)</td>
<td>1.07 (1.04, 1.10)</td>
</tr>
<tr>
<td>3</td>
<td>3.75 (3.62, 3.87)</td>
<td>-0.37 (-0.38, -0.35)</td>
<td>50.1 (50.0, 50.2)</td>
<td>1.16 (1.13, 1.19)</td>
</tr>
<tr>
<td>4</td>
<td>3.54 (3.44, 3.64)</td>
<td>-0.36 (-0.37, -0.34)</td>
<td>50.2 (50.1, 50.3)</td>
<td>1.25 (1.22, 1.28)</td>
</tr>
<tr>
<td>5-Most deprived</td>
<td>3.35 (3.27, 3.43)</td>
<td>-0.35 (-0.36, -0.33)</td>
<td>50.3 (50.1, 50.4)</td>
<td>1.40 (1.37, 1.43)</td>
</tr>
<tr>
<td><strong>IMD excluding health</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1-Least deprived</td>
<td>3.96 (3.81, 4.11)</td>
<td>-0.45 (-0.47, -0.44)</td>
<td>49.4 (49.2, 49.5)</td>
<td>Reference</td>
</tr>
<tr>
<td>2</td>
<td>3.85 (3.71, 3.98)</td>
<td>-0.42 (-0.43, -0.40)</td>
<td>49.7 (49.6, 49.8)</td>
<td>1.07 (1.04, 1.10)</td>
</tr>
<tr>
<td>3</td>
<td>3.71 (3.59, 3.83)</td>
<td>-0.37 (-0.38, -0.35)</td>
<td>50.1 (50.0, 50.2)</td>
<td>1.14 (1.11, 1.17)</td>
</tr>
<tr>
<td>4</td>
<td>3.60 (3.50, 3.70)</td>
<td>-0.36 (-0.37, -0.34)</td>
<td>50.2 (50.1, 50.3)</td>
<td>1.23 (1.20, 1.26)</td>
</tr>
<tr>
<td>5-Most deprived</td>
<td>3.33 (3.25, 3.41)</td>
<td>-0.35 (-0.36, -0.33)</td>
<td>50.3 (50.1, 50.4)</td>
<td>1.37 (1.33, 1.40)</td>
</tr>
</tbody>
</table>

*amplitude exponentiated
5.4. Discussion

This study adds to the evidence of the burden of bronchiolitis presented in Chapter 4, showing that in December in England between 2012 and 2016 40% of all infant hospital admissions included a bronchiolitis diagnosis. There is a clear socioeconomic gradient to these admissions, and at week 50 (peak admission week) infants born into the highest level of socioeconomic deprivation had a risk of admission 40% greater than the lowest group. The results suggest that the association between seasonality and bronchiolitis admission rates is marginally moderated by level of socioeconomic position, with increasing socioeconomic deprivation associated with less seasonal variation and a slightly delayed epidemic peak.

5.4.1. Strengths and weaknesses

The study data were derived from a national hospital administrative dataset, which enabled me to create a representative birth cohort with minimum selection bias and follow patients up over time. I was able to link the majority of infants to maternal records, which was crucial for acquiring socioeconomic data as this information was missing for infant records before 2014 (but available in maternal records). However, the probabilistic method used is likely to have included erroneous links, introducing some bias into the dataset. Using the clinical diagnosis...
of bronchiolitis without data on laboratory test results for specific viruses meant that I could not attribute the bronchiolitis admissions to particular pathogens, although previous work from England shows that 80% of bronchiolitis admissions in infants (defined using ICD-10 code J21) are due to RSV. More broadly, as in Chapter 4, this study is a measure of hospital use and therefore does not measure community or primary care burden of bronchiolitis.

I added harmonic functions to the regression model to account for sinusoidal patterns in a highly seasonal infection; however, the model underestimates the size of the amplitude across socioeconomic groups. Improved estimation could have been achieved with the addition of more harmonic pairs or by using splines; however, as mentioned in Chapter 4, these methods would have added undue complexity to the estimation of seasonal parameters required by my research aim. The likeness in the relative difference between amplitudes by IMD across observed and fitted rates (Figures 5.1 and 5.2) gives me confidence in the results.

5.4.2. Research findings: comparison with other studies

Objective 5A: the proportion of infant hospital admissions attributable to bronchiolitis

Overall, between 2012 and 2016, more than 1 in 7 of all hospital admissions in England among infants included a diagnosis of bronchiolitis. During December, this figure rose to 1 in every 2.5 admissions, highlighting the substantial toll this condition takes on hospitals in England during winter. In absolute numbers, infants from the lowest fifth of deprivation made up 33% of the 193,793 hospital admissions with a diagnosis of bronchiolitis between 2012 and 2016. In comparison, 13% of admissions were among children in the highest fifth.

It is important to note that more children in the general population are in more deprived groups; therefore, this measure may be less illustrative than relative measures (that take into account the overall size of the groups). When considered as a proportion of all admissions within the group, the proportions are far less stark; bronchiolitis comprised 16.3% of all admission in the most deprived group compared to 14.0% in the least deprived group. Taking into account the time spent at risk of hospital admission, there was a 1.5-fold difference between the most and least deprived groups (59.8 versus 41.0 per 1000 infant-years).
Objective 5B: the extent to which seasonality of bronchiolitis admissions is modified by socioeconomic position

This work presents a small, but arguably inconsequential, difference in epidemic timing—the two most extreme IMD groups had a 0.9 week difference in predicted peak—and no difference in the relative duration of the epidemic. Research from Western Australia found no difference in the seasonality of RSV positive specimens among aboriginal children (a population that experience high levels of socioeconomic deprivation) compared to non-aboriginal children. USA-based research, on the other hand, found a negative association between the proportion of the population from a black ethnic group (correlated with socioeconomic position of the population) and seasonal peak timing at the ZIP-code level.

Less seasonal variation in lower socioeconomic groups reflects the continued (but low level) admissions for bronchiolitis during warmer months of the year amongst these infants. My study found a greater relative difference in the rates between infants at the lowest point of the epidemic curve (mid-June), where the predicted admission rate was 10.2 per 1000 infant-years in the most deprived compared to 5.2 per 1000 infant-years in the least deprived IMD quintile. This may reflect an increased divide between risk factors for RSV transmission during summer months (such as time spent indoors and overcrowding) in combination with the influence of non-RSV pathogens at this time. Understanding precise viral aetiology of bronchiolitis by time of the year and socioeconomic group may help to delineate seasonal differences further. A study in New Zealand found that non-RSV bronchiolitis was more common in infants from areas of higher deprivation. Future work would benefit from linkage to surveillance datasets, such as Public Health England’s Second Generation Surveillance or Respiratory Datamart Systems, to assess these factors.

5.4.3. Implications of study findings

In the context of current UK guidelines for administering Palivizumab, the results present little evidence to support differential timing of interventions for bronchiolitis amongst socioeconomic groups in England at the current time. The work does, however, present substantial difference in admission rates across population subgroups irrespective of season. This study highlights that, at the very least, infants from poorer backgrounds should be considered a priority group for future interventions. Based on previous research, I hypothesise that a combination of risk factors, such as tobacco smoke exposure, premature birth, congenital heart and chronic lung disease, outdoor air pollution and damp, overcrowded housing conditions, contribute to these inequities. Further work using formal methods to establish pathways through which
these factors affect the risk of bronchiolitis is needed to guide the most appropriate interventions.

5.5. Chapter summary

In this chapter, I have highlighted the substantial burden caused by bronchiolitis on English hospitals in winter. This is particularly apparent in December when, on average between 2012 and 2016, almost 40% of all infant admissions included a diagnosis of bronchiolitis. Using harmonic analysis, I found only a marginal moderation in the seasonality of these admissions by level of socioeconomic position. The results suggest that differential timing of interventions may not be necessary for different socioeconomic groups, but, with a social gradient in the volume of admissions, highlight the continued inequity in admissions rates for bronchiolitis more broadly. Moving forward, investigating the precise viral aetiology of bronchiolitis by season and socioeconomic group, as well as the risk factors mediating the link between socioeconomic deprivation and bronchiolitis rates, will help to further unpick the reasons for inequities in rates of hospital admissions for bronchiolitis in the UK.
Chapter 6. Socioeconomic position and bronchiolitis—mediation through maternal smoking

Recap and chapter overview

In chapters 4 and 5, I explored the seasonality of annual bronchiolitis epidemics across England, and associated risk factors for bronchiolitis focussing on socioeconomic deprivation. This work showed a clear socioeconomic gradient to the rates of hospital admissions for bronchiolitis amongst infants in England. Using a dataset constructed from Scottish administrative data, this chapter will look more closely at one modifiable risk factor that sits on the pathway between socioeconomic deprivation and bronchiolitis admissions, answering thesis objective 2:

To establish to what extent maternal smoking mediates the association between early life socioeconomic position and bronchiolitis admission rates in infancy.

I discuss the causal inference framework used to guide the mediation analysis, and employ a novel method to examine plausible causal assumptions of my study. I then discuss the available mediation estimands, before choosing and applying the most appropriate method to answer my research question.

6.1. Background

A social gradient is present in the incidence rates of hospitalisation for bronchiolitis. My research presented in Chapter 5 shows that children born in England between 2011 and 2016 into the most deprived quintile of IMD had an admission rate of 62.0 per 1000 child-years (95% CI 61.5 to 62.5) compared to 41.1 per 1000 child-years (95% CI 40.5 to 41.7) among children in the least deprived quintile. I hypothesised that a combination of risk factors including tobacco smoke exposure, premature birth, congenital heart and chronic lung disease, outdoor air pollution and damp, overcrowded housing conditions, contribute to these inequities. This hypothesis is based on pre-existing research where the association between socially patterned risk factors and bronchiolitis admission rates has been examined. However, whilst these associational studies help to describe patterns of bronchiolitis by infant characteristics, they do not break down the pathways through which socioeconomic circumstances influence risk of infection.
Understanding the mechanisms through which an exposure is linked to an outcome is an important component of epidemiological research. Specifically, quantifying the relative magnitude of the impact of intermediate variables could guide options amenable to intervention (as well as the best timing for those interventions).\textsuperscript{368} This is particularly helpful in instances where intervening on the exposure is less feasible (as is arguably the case with socioeconomic position).\textsuperscript{369} However, traditional formulations of mediation analyses, which used product of coefficients and difference in coefficients estimators,\textsuperscript{370,371} have been shown to rely on very strong parametric assumptions and cannot be generalised beyond simple linear regression models.\textsuperscript{372,373} The latter are only suitable for settings where all relationships are linear and there are no exposure-mediator interactions, nor non-linearities or multiple interacting mediators.\textsuperscript{374} In many applied settings, the focus is on more than one mediator, either because the mediating mechanisms of interest are complex, or because the mediator of interest is influenced by earlier mediators (an example of mediator-outcome intermediate confounding).\textsuperscript{375} These methodological challenges may explain the lack of research on the complex pathways between socioeconomic position and bronchiolitis.

Advances in the causal inference literature, which draws on the counterfactual framework, has produced some solutions to the problem of decomposing the effects of mediators in complex settings.\textsuperscript{370} Several causal mediation estimands are now available to researchers—their choice dependent on the research question at hand and the plausibility of the required assumptions.\textsuperscript{376} As I will elucidate in this chapter, implementation of mediation analysis (even when focussing on a single mediator) is complex, requiring careful consideration of the causal structure and associated assumptions of the examined pathways.\textsuperscript{371,377} There are further intricacies specific to my study because of the complexities associated with defining the exposure, socioeconomic position, and the number of mediator-outcome confounders that are associated with this exposure.\textsuperscript{370} Drawing upon the causal inference framework, the aim of this chapter is:

To establish the extent to which maternal smoking during pregnancy mediates the association between parental socioeconomic position and bronchiolitis hospital admission rates in infancy.

I am focussing on maternal smoking because it is an established risk factor for bronchiolitis that is strongly socioeconomically patterned and, in comparison to other potential mediators such as birthweight and gestational age, presents a pathway that has practical implications in terms of interventions.\textsuperscript{131,138} There are 4 objectives underpinning this aim:

**Objective 6A** Outline the basic principles and assumptions of causal inference.
Objective 6B Use an evidence based approach to create a causal diagram to inform the analyses.

Objective 6C Choose the appropriate mediation estimand(s) with consideration to both the complexities of the process and the available data.

Objective 6D Based on findings from objective 6A-6C, establish the extent to which maternal smoking explains the socioeconomic disparity in hospital admission for bronchiolitis in infancy.

6.2. Causal inference framework

6.2.1. Basic principles

Typically, observational studies have been used to look at associations—showing the strength of relationship between two or more variables. The previous two studies in this thesis, for example, were concerned with the association between the time of year, population factors and hospital admissions for bronchiolitis. This type of analysis enabled me to describe the patterns, and estimate the risk, of bronchiolitis events across England under specified conditions. Judea Pearl writes that standard statistical analysis, which uses samples drawn from a population to estimate parameters, allows researchers to infer associations among variables where conditions remain constant. However, the same approaches cannot be applied for situations where we wish to infer risks under changing conditions, such as when an intervention is or is not implemented. To understand risks under these conditions, the focus needs to shift to understanding why the associations exist or, to put it another way, determining that a cause leads to an effect (i.e. causal inference).

Causal interpretation of observational associations can be unconvincing because, unlike in randomised trials, the exposure (or treatment) is not randomly assigned to individuals. Another way of putting this, is that association is largely not taken to mean causation in observational studies because the exposed and unexposed are not generally exchangeable. This is unlike randomised experiments, where randomisation is applied to ensure the exposed and unexposed are exchangeable. This means that the outcomes observed in one of the randomised groups can be taken to represent the outcomes from the other group if they had received the same exposure/treatment. However, as written by Pearl, the move from association to causation is achievable by including information provided through causal assumptions that “identify relationships that remain invariant when external conditions change.” Understanding these causal assumptions, detailed below, can be aided by the use of counterfactual reasoning.
Using counterfactual statements, research questions can be reframed to draw contrasts between outcomes given different sets of conditions. In this chapter, I adopt the potential outcomes framework to define causal parameters of interest (causal estimands),\textsuperscript{370,382–384} e.g. the total causal effect (TCE) on $Y$ (the outcome) of a change in $A$ (the exposure) from $a^*$ to $a$ is defined as $\text{TCE} = \mathbb{E}[Y(a) - Y(a^*)]$, where $\mathbb{E}$ stands for expectation in the population of interest, and $Y(a)$ is the potential outcome had the exposure $A$ been set to take the value $a$, possibly contrary to the fact, and $Y(a^*)$ is the potential outcome had exposure $A$ been set to take the value $a^*$, where $a \neq a^*$. More specifically, ‘what would the risk of bronchiolitis be if all infants were born into low versus high levels of socioeconomic deprivation?’ would be expressed as the TCE where $a^*$ and $a$ stand for either low or high socioeconomic deprivation. This type of reasoning can help researchers to consider the effects of an intervention without it being enacted, and highlight where limitations in reasoning and methods exist.\textsuperscript{385,386}

6.2.2. Key assumptions

Hernán and Robins describe conditions that, if met, may informally allow an observational study to be conceptualised as a conditionally randomized experiment.\textsuperscript{380} I will now define four assumptions, labelled no interference, consistency, positivity and exchangeability, that are usually invoked to derive causal effects with application to the current study.

No interference

No interference assumes that an individual’s potential outcome does not depend on other individuals’ exposure values.\textsuperscript{380,387} The presence of interference would mean that the counterfactual outcomes have more complex definitions and, thus, causal inference also becomes more complex.\textsuperscript{387} In my study, interference would be present if the risk of being admitted to hospital for bronchiolitis for one infant was dependent on the SEP of other infants. As shown in Chapters 3 and 4, there is a high concentration of hospital admissions for this condition in winter, driven by the seasonality of RSV. Children from lower SEPs do make up a higher proportion of those admitted; however, the continual rise in hospital admission rates for bronchiolitis over time suggests that infants with bronchiolitis are not turned away due to capacity issues. Al-Mahtot and colleagues, for example, showed that the odds of an emergency admission for hospital in Scotland with a diagnosis of bronchiolitis was 24% greater in 2010-13 relative to 2000-03.\textsuperscript{354} It is suggested that duration of stay may be influenced by bed capacity;\textsuperscript{327} however, this is not the outcome I am looking at in my study. For these reasons, I assume that the assumption of no interference is upheld in my study.
Consistency

Under the consistency assumption, write Hernán and Robins, “the values of treatment under comparison correspond to well-defined interventions that, in turn, correspond to the versions of treatment in the data”. Put another way, the exposure definition must have enough precision that any variation in that exposure does not lead to a different outcome. There are two components to consider here in relation to my exposure: whether SEP can actually be intervened upon; and, if so, whether it is defined well enough to meet the consistency assumption. As discussed at the beginning of Chapter 1, intervening on SEP is not an impossible goal, but neither is it an easy one; requiring major and sustained social change.

Even after putting the argument about intervening directly upon SEP aside, however, Rehkopf et al. argue that composite measures (such as the Scottish IMD, used in this study) are not well defined interventions and, for this reason, violate the consistency assumption. They recommend avoiding composite measures altogether. Other non-composite measures of SEP such as education or income lend themselves better to the constraints of the consistency assumption. However, as explained from the outset of this thesis, IMD is advantageous in the comprehensive and nationally relevant measurement of inequity it provides.

By focussing on the mechanisms underpinning inequities, there are ways to circumvent the strong requirements needed to meet the consistency assumption. Naimi and colleagues show how mediation methods can be used to consider how upstream risk factors for health are explained by proximal risk factors by allowing the statistical association between exposure and outcome to carry “relevant disparity-related (but not necessarily causal) information”. This is the approach I will adopt in my study and is discussed in further detail in Section 6.4.2.

The positivity assumption

Positivity means that all individuals have a probability greater than 0 (a positive probability) of being assigned all values of the exposure variable, in every stratum defined by the covariates C that allow for the next assumption to be met, i.e. the confounders. In the current study, there must be infants from low and high socioeconomic groups in each stratum of the confounder variables to meet this assumption. Given the large size of the birth cohort to be used in my study, it is likely that this assumption will be met; however, this may be dependent on the number and rarity of confounders. I will revisit this assumption after these aspects have been examined in my dataset.

The exchangeability assumption
Exchangeability, as applied in the current study, assumes that infants born into high and low socioeconomic position groups (i.e. the unexposed and exposed) are comparable in every respect apart from the exposure. If this is not the case, then it cannot be expected that the observed outcome (hospital admission for bronchiolitis) for the exposed is the same as the potential outcome under exposure of the unexposed. As exchangeability is unlikely to occur in observational data, we can instead aim for achieving conditional exchangeability in order to elicit a valid causal interpretation, conditional on a set of covariates $C$. Thus, we assume that there are no uncontrolled confounding variables, which means that within strata defined by these confounders, the exchangeability condition that is guaranteed in randomised controlled trials is met. Hernán and Robins describe this as “the conditional probability of receiving every value of treatment, though not decided by the investigators, depends only on measured covariates $[C]$”. Causal diagrams (introduced next, Section 6.3) helps to identify the confounders necessary to condition on to achieve conditional exchangeability.

### 6.3. Causal diagrams

The identification of the set of confounding variables $C$ needed to meet the conditional exchangeability assumption can be aided by the use of directed acyclic graphs (DAGs). DAGs are a simple method of graphically representing the assumptions regarding how the variables in a study and their common precursors are causally related to one another. DAGs consist of nodes (representing variables) and unidirectional edges (or arrows, representing potential causal effects) that are either present or missing between nodes. Causal DAGs are DAGs that include all common causes of any nodes in the diagram and are drawn starting from the main exposure-outcome relationship. Causal DAGs can be used as a tool to illustrate both the assumptions made in causal analysis and the wider context of the causal questions being asked. Such diagrams help to identify: whether there is confounding in the exposure-outcome relationship and which variables need to be controlled for, or not controlled for, when estimating the effects of interest. A missing edge represents the strongest assumption in a DAG: that there is no direct causal effect between two variables.

There is a lack of best practice guiding the specification of DAGs and, where studies have used them, the diagrams have been criticised for being overly simplistic and altered to fit available data. Further, DAGs aren’t always presented in studies that claim to use them. For these reasons, Ferguson and colleagues created an evidence synthesis for constructing DAGs (ESC-DAG). The ESC-DAG primarily serves to guide DAG development from background knowledge and direct primary data analysis, whilst examining plausible causal assumptions.
Evidence synthesis for constructing directed acyclic graphs (ESC-DAG) application

To apply the ESC-DAG to my research question I started by conducting a search of the relevant literature. An initial search for studies that had looked at all three components of my research question (socioeconomic position, smoking and bronchiolitis) led to just one result, which was a study that looked at the second-hand smoking after birth. I therefore broadened my search question to ‘how does early life socioeconomic position influence rates of bronchiolitis in early childhood?’ (Table 6.1). This enabled me to include more studies and consider other pathways beyond smoking, which is necessary to deal with the potential confounding of the relationship between the mediator (maternal smoking) and the outcome (bronchiolitis admissions).

Table 6.1. Search strategy for literature used in ESC-DAG

<table>
<thead>
<tr>
<th>Research question</th>
<th>How does early life socioeconomic position influence rates of bronchiolitis in early childhood?</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Search strategy</strong></td>
<td></td>
</tr>
<tr>
<td><strong>Databases</strong></td>
<td>Medline, Embase, Maternity and Infant care database (using OvidSP) and PubMed</td>
</tr>
<tr>
<td><strong>Search terms</strong></td>
<td>OvidSP: (Social determinants of health/ OR Social status/ OR Social structure/ or Socioeconomics/) AND (bronchiolitis OR human respiratory syncytial virus/). PubMed: (Socioeconomic factors [MeSH term] OR Social class [MeSH term]) AND (bronchiolitis OR human respiratory syncytial virus)</td>
</tr>
</tbody>
</table>
| **Inclusion criteria** | • English language  
                        | • Human                                                      |
|                    | • High-income countries                                                                |
|                    | • Study participants <2 years                                                           |
|                    | • Published ≥2000                                                                      |
|                    | • Outcome is bronchiolitis cases or rates                                             |
|                    | • Longitudinal, cohort or prospective research                                         |
| **Identified papers** |                                                                                     |
| **Initial number** | 72 studies                                                                            |
| **Removed**        | 17 duplicates                                                                         |
|                    | 46 not relevant (from title/abstract)                                                  |
|                    | 73 not relevant (from paper)                                                           |
| **Final number**   | 9 studies                                                                              |

Using the list of identified and relevant studies, ordered descending by year of publication, I carried out the 3-stage process outlined below. The process closely follows the ESC-DAG protocol described by Ferguson and colleagues, with some adjustments that are explicitly noted in the text below:
**Stage 1.** Firstly, I noted the main aspects of the study, including the study design, statistical methods, study exposure(s), study outcome(s) and control variables, in a decision log (Table A6.1, Appendix 6.1). I used this information to map the conclusions of each study to an initial graph called an implied graph. For each implied graph, the following process was followed: a directed edge was drawn from the exposure to the outcome; all control variables were entered into the graph; mediators were mapped out based on the study conclusions; and, finally, the implied graph was saturated by drawing edges between all nodes. To reduce the high volume of assessments required, I excluded variables at this initial stage that were not applicable to a UK context (and included a reason for the exclusion in the decision log). For example, presence/type of health insurance was used in Inagaki et al.’s study of bronchiolitis in New York State as one of two indicators of SEP, but is less applicable in the context of the publicly funded NHS in the UK.

**Stage 2.** The second stage, translation, involved applying causal theory to each relationship in the implied graph, including consideration of temporality, face-validity and prior theory/research. I then applied a counterfactual thought experiment; hypothesising the potential outcome that would occur if all individuals were set to receive the same counterfactual exposure (for example, if all infants were born female). At this stage, a directed edge could be retained, reversed or removed. Importantly, it is a stronger assumption to have a line removed then to let it remain. Again, the theory/evidence was considered in relation to the UK rather than the country where the study was conducted. Each step of the translation was recorded in an Excel spreadsheet (not shown) and summarised in the resultant DAG for each assessed study (Table A6.1, Appendix 6.1). Relationships that had already been assessed in other studies were not re-assessed.

**Stage 3.** When all papers were assessed, a final index of all directed edges was created (Table 6.2). Similar nodes were combined at this stage to avoid repetition, as well as for purposes of parsimony. For example various socioeconomic measures, including household income, neighbourhood SEP and social exclusion risk, were combined into the overarching parental SEP. This index was used to create the integrated-DAG representing the relationship between parental SEP and infant hospital admission for bronchiolitis, shown in Figure 6.1, below. I will use the integrated-DAG to guide confounder selection in the mediation analysis in section 6.5 in this chapter and again in Chapter 7.

DAGitty, an online open source graphical tool, was used to create the DAGs presented in Table A6.1 and Figure 6.1, and to determine minimally sufficient adjusted sets for the main analyses.
Table 6.2. Final directed edge index

<table>
<thead>
<tr>
<th>#</th>
<th>Edge originates from</th>
<th>Edge terminates at</th>
<th>#</th>
<th>Edge originates from</th>
<th>Edge terminates at</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Parental SEP</td>
<td>Bronchiolitis</td>
<td>76</td>
<td>Rural/urban</td>
<td>Bronch. dysplasia*</td>
</tr>
<tr>
<td>2</td>
<td>Child ethnicity</td>
<td>Bronchiolitis</td>
<td>77</td>
<td>Congenital anomaly</td>
<td>Bronchiolitis</td>
</tr>
<tr>
<td>3</td>
<td>Sex</td>
<td>Bronchiolitis</td>
<td>80</td>
<td>Parental SEP</td>
<td>Congenital anomaly</td>
</tr>
<tr>
<td>4</td>
<td>Delivery method</td>
<td>Bronchiolitis</td>
<td>81</td>
<td>Sex</td>
<td>Congenital anomaly</td>
</tr>
<tr>
<td>5</td>
<td>Child birth season</td>
<td>Bronchiolitis</td>
<td>82</td>
<td>Congenital anomaly</td>
<td>Gestational age</td>
</tr>
<tr>
<td>6</td>
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<td>Bronchiolitis</td>
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*bronch. dysplasia = bronchopulmonary dysplasia*
DAGitty identified the minimally sufficient adjustment set for estimating the direct effect of parental SEP on hospital admission for bronchiolitis as:

- area of residence,
- birth year,
- maternal country of birth
- and maternal ethnicity

These are indicated by red nodes in the diagram.

The yellow node is the exposure, the blue node with a black outline is the outcome and all other blue nodes are ancestors of the outcome (and not confounders of the exposure-outcome relationship).

Figure 6.1. Overall integrated-DAG created through the ESC-DAG protocol*
6.4. Causal mediation analysis

6.4.1. Controlled, natural and interventional (in)direct effects

There are different approaches to study mediation in the causal literature, some of which attempt to separate the total effect of an exposure on an outcome into effects that act directly from the exposure or via a particular mediator (or set of mediators), respectively, the direct and indirect effects. In my study, the indirect effect of interest is the effect of socioeconomic position on the risk of hospital admission for bronchiolitis that involves maternal smoking. I will now present the main approaches to mediation analysis, discussing the applicability of each to my research question.

Controlled direct effects

Controlled direct effects (CDEs) quantify the effect of the exposure when the mediator is set to a specific value for everyone in the population. The CDE of \( A \) on \( Y \), not mediated via \( M \) when expressed as a linear contrast, is defined as \( \text{CDE}(m) = \mathbb{E}[Y(a,m) - Y(a^*,m)] \), where \( Y(a,m) \) is the potential outcome when \( A \) is set to take the value \( a \) and \( M \) the value \( m \). This is the contrast between the potential outcomes under exposure \( A=a \) versus \( A=a^* \), where the mediator is set at the same fixed level for everyone in the population (\( M=m \)). The CDE(m) may vary with values of \( m \). Moreover the CDE could be expressed on different scales, for example as ratios of expected potential outcomes (when the outcome is binary).

CDEs are described as relevant to interventional settings where policy evaluation is of interest, in particular when the mediator is binary or categorical, such as maternal smoking in my study. However, it is difficult to ascertain effect decomposition with CDEs, with Naimi et al. describing the controlled indirect effect as “some contrast between the total and controlled direct effects in the absence of exposure-mediator interactions”. This is particularly problematic where more than one mediator is present. Alongside the two technical assumptions of no interference and consistency for the outcome, the two main identification assumptions required to ascertain CDEs are:

1. no uncontrolled (or unmeasured) confounding of the exposure-outcome relationship
2. no uncontrolled (or unmeasured) confounding of mediator-outcome relationship
Natural (in)direct effects

Another causal mediation strategy quantifies effects based on exposure and mediator values that naturally occur in the population. Pearl writes that these “natural effects” are a descriptive and more subtle interpretation of effects compared to the CDEs. The natural direct effect (NDE) of $A$ on $Y$, not mediated via $M$ (when expressed as a linear contrast), is defined as $\text{NDE} = E[Y(a,M(a*))] - E[Y(a*,M(a*)])$, where $M(a*)$ is the potential mediator had the exposure been set to take the baseline value $a^*$. This is the contrast between potential outcomes under exposure $A=a$ versus $A=a^*$, where the mediator is set to the value it would have taken under a reference (i.e. “natural”) exposure value ($M=a^*$). These values are specific to each individual, unlike the mediator value $m$ used in the CDE definition. The natural indirect effect (NIE) of $A$ on $Y$ that operates via the mediator $M$ is defined as $\text{NIE} = E[Y(a,M(a))] - E[Y(a,M(a*)])$. This is the contrast between the potential outcomes where $A$ is set to the same exposed value ($A=a$), but the mediator is set to take the value it would have taken had the exposure been set to exposure ($A=a$) or no exposure ($A=a^*$).

These definitions allow for effect decomposition, with the NDE and NIE summing to the TCE. Natural effects are seen as particularly appropriate for providing information on the actions of mechanisms. However, the identification assumptions required for natural effects are considerably stronger than for CDEs, making this method also unsuitable for my research question. Alongside the two conditionally exchangeability assumptions listed for CDEs (no uncontrolled confounding of the exposure-outcome nor the mediator-outcome relationships), and consistency for mediator and outcome, estimating natural effects additionally requires that:

3. there is no uncontrolled confounding of the exposure-mediator relationship,
4. the potential outcome under exposure and set mediator value, $Y(a,m)$, and the potential mediator under no exposure, $M(a^*)$, need be independent conditionally on the baseline founders.

This fourth assumption is known as the “cross-world independence assumption” because it leads us to simultaneously contemplate worlds where an individual is exposed (the potential outcome under exposure) and unexposed (the potential mediator under no exposure). In my study, this would require imagining an infant born to parents with high socioeconomic deprivation and, at the same time, the same infant born to parents with low socioeconomic deprivation. As this “recanting witness” exposure assignment could not occur in real life, the reliance of causal mediation analysis on this assumption has been criticised as being empirically unverifiable. This assumption however implies that there are no variables that
are downstream from the exposure and that confound the mediator-outcome relationship. This may be a defensible assumption in certain settings where there is a small time-gap from exposure to mediator.\textsuperscript{402} For example, if an educational video about smoking (exposure) led to a quick change in attitude (mediator), which, in turn, led to smoking cessation (outcome). In our application, however, this assumption is less defensible. Indeed, it is argued that this assumption diminishes the public health relevance of the NDE and NIE.\textsuperscript{403}

Randomised interventional analogues of the NDE and NIE

Alternative mediation estimands have been introduced that can be identified under weaker assumptions than the NDE and NIE, for example, the randomised interventional analogues (RIA) of the NDE and NIE.\textsuperscript{384} The definitions for these estimands are the same as for the NDE/NIE, above, but differ in respect to the values the mediators are set. RIA-NDE=$E[Y(a,M_m^*)-Y(a^*,M_m^*)]$, a quantity that captures the direct effect of $A$ on $Y$ not involving $M$, when the contrast is between the potential outcomes under exposure $A=a$ versus $A=a^*$, where the mediator is set to take a value that is randomly drawn from its conditional counterfactual distribution under no exposure, conditional on the confounders $C$. It is interpreted as the effect of $A$ that would remain if the distribution of the mediator $M$ could be changed to be the same as the distribution of the mediator had $A$ been set to $a^*$. RIA-NIE=$E[Y(a,M_c)-Y(a,M_c^*)]$, captures the effect of $A$ on $Y$ operating via the mediator $M$, where the value of the mediator is set to take a value randomly drawn from its conditional counterfactual distribution under $A=a$ versus a value drawn from its conditional counterfactual distribution under $A=a^*$, while the exposure is set to remain fixed at the same value $a$.

I present an example where these mediational effects are estimated in the presence of a single mediator, gestational age, in Box 6.1.\textsuperscript{384} I estimated the RIA-NIE of gestational age as the counterfactual difference in bronchiolitis admission rates between children born in the high versus low deprivation group. Here, gestational age (in weeks) is drawn at random from the distribution of gestational age among infants in the low socioeconomic deprivation group, conditionally on confounders, whilst levels of socioeconomic deprivation remained high. The effects do not lead to a decomposition of the TCE but, as reflected by Daniel and De Stavola, there is trade-off to be made between “how closely an estimand corresponds to what we truly wish to learn about, and the strength of the assumptions needed to identify it”.\textsuperscript{373 p15} Despite not providing insight into all pathways, this method is arguably more applicable to real world and policy-relevant scenarios.\textsuperscript{406}
Chapter 6 - Mediation through maternal smoking

Box 6.1. Could socioeconomic inequity in bronchiolitis admissions be reduced through intervening on gestational age?*

**Aim** To establish the extent to which socioeconomic inequity in hospital admissions for bronchiolitis would be reduced if socioeconomic disparities in gestational age were eliminated. In this study, socioeconomic inequity refers to the difference in outcomes between infants born to parents in high versus low levels of socioeconomic deprivation.

**Methods** Children born in English NHS hospitals between 2011 and 2016 were followed up for one year using HES APC records (N=3,948,819). The exposure, socioeconomic deprivation, was measured using the Index of Multiple Deprivation 2010 score assigned to the infant's registered lower super output area at birth (split into high or low levels of deprivation). Gestational age (in weeks) and sex (male or female) were also retrieved from hospital records at birth. Presence of congenital anomalies (yes/no), based on the Hardelid UK chronic condition ICD-10 code list, were identified in infant hospital admissions or death certificate up to age two. The study outcome was any emergency admissions to hospital with a diagnoses of bronchiolitis in the first year of life (identified in hospital records by ICD-10 code J21). I estimated the RIA-NIE of gestational age using g-computation (as implemented in Stata using the gformula command), while controlling for the confounders of the mediator and outcome association—congenital abnormalities and sex. I also estimated the total causal effect (TCE) of socioeconomic deprivation on hospital admission rates for bronchiolitis, controlling for the same confounders.

**Findings** On average, between 2012 to 2016, the bronchiolitis admission rate was 41.0 per 1000 infant-years (95% CI 40.6 to 41.4) among infants in the low deprivation group compared to 51.6 per 1000 infant-years (95% CI 51.2 to 52.0) in the high deprived group. Assuming no unmeasured confounding, the estimated TCE indicates that, if socioeconomic disparities could be removed, the rate of admissions overall would decrease by 9.68 per 1000 infant-years (95% CI 9.17 to 10.18). The estimated RIA-NIE indicates that if socioeconomic disparity remained but the distribution of gestational age were set to that of the low deprivation group, the rate of admissions would decrease by 1.40 per 1000 infant-years (95% CI 1.31 to 1.42). This is the equivalent to removing 14.0% (95% CI 13.2 to 14.8) of the difference in admission rates due to socioeconomic deprivation.

*An earlier version of this abstract was published as a meeting abstract in the Lancet (included in Appendix 6.2).*
6.4.2. Multiple mediators

The methods and example presented above consider only a single mediator. However, as Steen and colleagues argue, even where researchers are interested in studying a single mechanism, the presence of other mediators cannot simply be ignored. Vanderweele describes an informal approach to multiple mediators whereby mediators are assessed one at a time and the proportion mediated summed to get the total indirect effect. However, for this method to produce reliable estimates, mediators must be conditionally independent given the exposure and baseline confounders. As shown in Figure 6.1, mediators on the pathway between SEP and bronchiolitis, such as maternal asthma and maternal age, are not causally independent from maternal smoking, even after controlling for possible confounders. The conditional independence assumption is therefore broken, which means that I cannot simply replicate the analysis in Box 6.1 with alternative mediators using the RIA approach. This would likely produce estimates that together do not meaningfully correspond to the total association.

A potential solution to the complex issue of intermediate confounding lies with counterfactual disparity measures (CDMs). By setting the mediator to a predefined value (e.g. \( m=0 \)), CDM captures the proportion of outcome disparity due to the exposure that would remain if a mediator (i.e. maternal smoking) were intervened upon and set to a chosen value \( m \), without intervening on \( A \). Formally defined, \( \text{CDM}(m) = E[Y(m)|A=1] - E[Y(m)|A=0] \), where \( Y(m) \) is the potential outcome had the mediator \( M \) been set to take the value \( m \), and where \( |A=1| \) and \( |A=0| \) indicate respectively that the expectation is in the exposed/unexposed subpopulation.

CDMs have two major benefits: they can still be identified in cases where there are confounders of the mediator-outcome relationship that are on the causal pathway from the exposure (such as maternal asthma and maternal age), as long as they are controlled for; and the identifiable assumption of no uncontrolled confounding of the exposure-outcome and exposure-mediator relationships are not required. This is because we wish only to characterise the overall disparity with this measure and therefore we are not concerned with whether the exposure-outcome association is confounded. The assumption of no unmeasured mediator-outcome confounding still applies. Although confounders of the exposure-outcome relationship are not required when estimating the CDMs, they can be included produce comparisons adjusted for these variables.

To address my aim, and given the difficulties in meeting the assumptions to be invoked to estimate natural effects (which would allow me to establish the extent to which maternal smoking mediates the association between early life socioeconomic position and bronchiolitis admission rates in infancy), I will instead focus on estimating the CDM of setting the mediator to be equal to non-smoking. This requires a reformulation of objective 6D, which becomes
“establish the magnitude of socioeconomic disparity in infant hospital admission for bronchiolitis that would remain if no women smoked during pregnancy”, defined as $CDM(m=0)=E[Y(m=0)|A=a]-E[Y(m=0)|A=a^*]$, where $A =$ socioeconomic deprivation (a if high/medium, a* if low), $M =$ maternal smoking (1 if yes, 0 if no) and $Y =$ hospital admission for bronchiolitis within the first year of life (1 is yes, 0 if no). This captures the contrast between the potential risks of hospital admission for bronchiolitis under high compared to low levels of socioeconomic deprivation, if all mothers were non-smokers.

In the following analyses, I will first identify, and then adjust for, confounders of the relationships between socioeconomic deprivation and smoking, and smoking and bronchiolitis cases.

### 6.5. Study cohort and variable selection

#### 6.5.1. Study DAG

I adapted Figure 6.1 to look at confounders of the exposure-outcome (parental SEP-bronchiolitis admission) and mediator-outcome (maternal smoking-bronchiolitis admission) relationships. The final study DAG is shown in Figure 6.2. To satisfy the assumption of no unmeasured mediator-outcome confounding, the minimal adjustment set is area of residence, maternal ethnicity (or child ethnicity), maternal country of birth, year, maternal age, maternal asthma, and parity. Confounders of the exposure-outcome relationship are area of residence, year, maternal ethnicity and maternal country of birth. Other covariates in the ESC-DAG (Figure 6.1) are not included here because they are all downstream from parental SEP and maternal smoking so are therefore not confounders in either the exposure-outcome or mediator-outcome relationships.
6.5.2. Data sources and study cohort

I used a subsection of the birth cohort of Scottish children described in Chapter 3 as the cohort for this study. I created this cohort using several linked administrative health datasets obtained via the Electronic Data Research and Innovation Service (eDRIS), who assigned a unique, pseudonymised identifier to each individual and deterministically linked children to their mothers via SMR-02. This allowed me to link records belonging to the same child and to mother-child pairs throughout the datasets. The linked datasets that formed the cohort are shown in Table 6.3.

Births are required by law to be registered with NRS within 21 days of the event and this register was used to form the core spine of the cohort.291 The birth registration data were supplemented by the mother's delivery record, which was retrieved from the SMR02, a dataset submitted by maternity hospitals on inpatient and day case activity relating to obstetrics. Information about deaths was retrieved from NRS death registrations and emigrations from
the CHI registry. Non-birth related hospital admissions in children were retrieved from SMR01, which includes information on inpatients and day cases discharged from hospitals in Scotland. SMR01 records include diagnostic, procedure and operation information, as well as non-clinical information such as location and length of stay. Data were also extracted from PIS, which contains information on medicines prescribed and dispensed within the community setting in Scotland (not including hospital administered/dispensed medication).

The cohort is defined by children born between 1st January 2012 and 31st December 2016 in Scotland. Their follow-up began at birth and continued until the child’s 1st birthday. Children born to mothers of non-Scottish residency (identified through mother’s country of residency in the NRS birth registrations) were excluded from the dataset to prevent potential systematic loss to follow up. Children had to be alive for at least one day to be included in the cohort. One child from each non-singleton birth was randomly selected to be included in the study.

Table 6.3. Scottish datasets used to create the study cohort for this study

<table>
<thead>
<tr>
<th>Acronym</th>
<th>Dataset</th>
<th>Information</th>
<th>Variables</th>
</tr>
</thead>
<tbody>
<tr>
<td>NRS</td>
<td>National Records of Scotland birth &amp; death registrations</td>
<td>Child</td>
<td>Postal area, SIMD, date of birth, date of death, mother country of birth, mother country of residence</td>
</tr>
<tr>
<td>CHI registry</td>
<td>Community Health Index registry</td>
<td>Child</td>
<td>Date of emigration</td>
</tr>
<tr>
<td>SMR02</td>
<td>Scottish Morbidity Records – obstetric related hospital admissions</td>
<td>Mother</td>
<td>Smoking status, age at delivery, parity, SIMD</td>
</tr>
<tr>
<td>SMR01</td>
<td>Scottish morbidity records–general hospital admissions</td>
<td>Child and mother</td>
<td>Diagnoses, date of admission, date of discharge</td>
</tr>
<tr>
<td>PIS</td>
<td>Scottish national prescribing information system</td>
<td>Mother</td>
<td>Medication name, date dispensed</td>
</tr>
</tbody>
</table>

6.5.3. Study variables

Outcome

The study outcome was any emergency hospital admission with a main or secondary diagnosis of bronchiolitis in the first year of life (yes/no). Bronchiolitis admissions were identified in SMR01 by the ICD-10 code J21 as a primary or any of the five secondary diagnosis codes.
Exposure

I used Scottish Index for Multiple Deprivation (SIMD) 2012 to capture socioeconomic position for each cohort member. SIMD is a relative measure of deprivation based on small areas called data zones (the Scottish equivalent of lower super output areas). SIMD combines scores of data zone-level deprivation across seven domains: income, employment, health, education, access to services, crime and housing. SIMD 2012 deciles (which are based on the data zone of residential addresses) were retrieved from NRS birth registration files and supplemented from SBR or SMR02 where missing. Deciles are constructed by ranking data zones by SIMD scores and dividing these into ten equal groups. SIMD deciles were split into three groups for analyses indicating high (top 30% rank of SIMD scores), medium (middle 40% rank of SIMD scores) and low levels of socioeconomic deprivation (bottom 30% rank of SIMD scores). Three groups as opposed to five were used because of the complications involved in estimating and interpreting results from the mediation analyses. I could not construct exact SIMD tertiles from the data because raw SIMD scores or data zones (to assign SIMD scores manually) were not available in my dataset.

Mediator

Mothers smoking status, stored in the SMR02, was recorded by midwives at first antenatal appointment (at around 8-12 weeks gestation). A binary variable, maternal smoking (yes/no), was created from this information with yes signifying that traditional cigarettes (i.e. non-e-cigarettes) were smoked at any point during their pregnancy and no for no smoking during pregnancy. E-cigarette smokers and women who smoked prior to their pregnancy were placed in the non-smoking category.

Confounders

Date of birth and postcode area were retrieved from each child’s birth registration file (NRS). There are 16 postcode areas in Scotland, which were grouped to make 5 regions with similar population sizes as shown in Figure 6.3. SMR02 provided mother’s age at delivery, which was split into four groups (≤19, 20-29, 30-39 and ≥40 years) and older siblings (dichotomised into yes or no). Mother’s country of birth also was retrieved from the NRS database and dichotomised into UK or non-UK born. Maternal asthma was defined using SMR01 hospital admission and PIS data. Mothers were recorded as having asthma during pregnancy (yes/no) if, in the year preceding their child’s birthdate, they had:
• ≥1 hospital admission that included a diagnosis of asthma (ICD-10 code J45, including all subcategories);
• and/or ≥2 asthma medications dispensed. Medication was defined using the British National Formulary (BNF) and included adrenoreceptor agonists (BNF codes 3.1.1), theophyllines (BNF codes 3.1.3), inhaled corticosteroids (BNF codes 3.2), cromoglicates and related therapies (BNF codes 3.3.1) and leukotriene antagonists (BNF codes 3.3.2).

Ethnicity was not available in infant birth records and maternal ethnicity was missing in SMR02 for >25% of the cohort so this variable was not included in the main analysis.
1. **Northern Scotland**
   - HS – Harris
   - IV – Inverness
   - KW – Wick
   - ZE – Shetland

2. **Central Scotland**
   - AB - Aberdeen
   - DD – Dundee
   - FK – Falkirk
   - KY – Kirkcaldy
   - PA – Paisley
   - PH - Perth

3. **G – Glasgow**

4. **EH – Edinburgh**

5. **Southern Scotland**
   - DG - Douglas
   - KA - Kilmarnock
   - ML - Motherwell
   - TD - Galashiels

---

**Figure 6.3.** Map of Scotland, by postal area; key showing 5 grouping used in this study
6.6. **Statistical methods**

Cohort characteristics and missing data were firstly described using descriptive statistics. The association between covariates and the probability of missingness in at least one study variable was explored using multivariable logistic regression. These findings were used to decide how to treat missing data in this study.

6.6.1. Admission rates

Replicating the procedure for the English birth cohort (Chapters 3 and 4), I calculated the incidence rates of bronchiolitis admissions per 1000 infant-years. All admissions (i.e. not just the first admission) with bronchiolitis recorded as either the primary or secondary diagnosis for an infant during their first year of life were included in the analyses. To calculate admission rates, any admission for bronchiolitis within 14 days of discharge from another bronchiolitis admission was assumed to be associated with the same infection, and therefore only the first of these admissions was included in the analyses. Children were followed from birth or 1st January 2012, whichever was later, to their 1st birthday, date of death, date of emigration or 31st December 2017, whichever came first. Incidence rates were calculated by dividing the number of admissions by person-time at risk and expressed as admission-based rates per 1000 infant-years for each of the key study variables.

6.6.2. Disparities in bronchiolitis admissions

I used two approaches to estimate the magnitude of socioeconomic disparity in hospital admission for bronchiolitis that would remain if smoking during pregnancy were eliminated (the \( CDM(m=0) \)). Namely, inverse probability weighting (IPW) of marginal structural models (MSMs) and g-estimation of structural nested models. Both are part of the family of methods known as Robins’ generalised methods or g-methods, which can provide estimates of contrasts of potential outcomes with less of the identification conditions required by standard regression models.\(^{411}\) Alongside the \( CDM(m=0) \), I quantified the marginal risk of bronchiolitis by SIMD group and the marginal risk difference between SIMD groups. The \( CDM(m=0) \) and risk difference were used to calculate the proportion of the socioeconomic disparity in bronchiolitis that can be explained by maternal smoking. I computed bootstrap confidence intervals for these estimates using 1000 bootstrap replications. Stata 15.0 was used for data analysis,\(^{314}\) the code for which can be found in Appendix 6.3.

In the following two approaches:
• the CDM($m=0$) estimates capture the potential risk differences between high and low SIMD group, and medium and low SIMD group, in ≥1 hospital admission with a diagnosis of bronchiolitis during infancy per 1,000 live births if no mothers smoked during pregnancy;
• confounders of the mediator-outcome relationship were year of birth, maternal age, maternal country of birth, maternal asthma and area of residence;
• and confounders of the exposure-outcome relationship were maternal country of birth, year of birth and area of residence (which were included to calculate average effects across these characteristics).

Inverse probability weighting of marginal structural models

IPW of MSMs can be used to estimate CDMs on the assumption that the mediator model is correctly specified as a function of all mediator-outcome confounders (in addition to assumptions of no interference, consistency, positivity and conditional exchangeability, see Section 6.2.2). To calculate the CDM($m=0$) using this approach, I firstly derived two sets of weights (see also Box 6.2). The first set of weights were the inverse of the predicted propensity from a multinomial logistic regression model for the (categorical) exposure, SIMD groups, adjusted for exposure-outcome confounders. The second set of weights were the inverse of the predicted propensity from a logistic model for the (binary) mediator, maternal smoking, adjusted for mediator-outcome confounders (and exposure-outcome confounders, although in this instance the confounders are already included as mediator-outcomes confounders). The two sets of weights were multiplied together to create the final inverse probability weights. I then fitted a weighted logistic regression model of the outcome on the exposure, the mediator and the interaction between the exposure and mediator using the inverse probability weights. The estimated coefficient for the mediator in this model represents the estimated CDM($m=0$).

**G-estimation of structural nested models**

This method specifies associational models for the outcome, the exposure and the mediator and is doubly robust against invalid inferences because estimators are consistent if either the outcome or the mediator model is correctly specified. G-estimation of structural nested models can be used to estimate CDM($m=0$) as these can be viewed as a component of structural nested models. The steps, adapted from Naimi et al., are outlined in Box 6.3. A logistic regression model including exposure-outcome and mediator-outcome confounders was used to predict the mediator and a multinomial logistic regression model including exposure-outcome confounders was used to predict the exposure.

**Marginal risk difference**
To estimate the marginal risk difference, I quantified the potential risk of at least one bronchiolitis admission per 1000 live births by SIMD group and took the absolute difference in these potential risks. I present differences between the high and low SIMD groups and high and medium SIMD groups. As with the CDM($m=0$), these quantities were estimated using two different approaches.

To complement the CDM($m=0$) estimated using the IPW of MSMs approach, a logistic regression model of the outcome against the exposure was estimated using IPW for the exposure via Stata’s `binreg` command (with the risk difference option specified). The weights were found by logistic regression modelling of the exposure that includes the exposure-outcome confounders. The estimated constant from this model can be interpreted as the marginal potential risk of bronchiolitis admission if everyone was assigned to the baseline (low deprivation) SIMD group and the estimated coefficient for each level of the exposure as the estimate of the relevant risk difference.

To complement the g-estimation of structural nested model method, logistic regression models of the outcome, adjusted for exposure-outcome confounders, were fitted for each level of the exposure and standardised across the cohort. The mean of the predicted estimates give the standardised potential risk of admission for bronchiolitis for each SIMD group. The risk differences are derived by taking one mean from another.

Proportion of inequity explained by maternal smoking

Using the estimates derived above for the CDM($m=0$) and marginal risk difference, I calculated the total proportion of the inequity between SIMD groups that is explained by maternal smoking as:

$$\text{disparity reduction (\%)} = \frac{(\text{risk difference}-\text{CDM}(m=0))}{\text{risk difference}} \times 100$$

Two estimates of the disparity reduction were calculated, one for each approach. The two approaches rely on different parametric assumptions (when using IPW that the propensity score models for the exposure and mediator are correctly specified, when using standardisation that the outcome and mediator models are correctly specified) and the use of both therefore allows me to assess the robustness of the estimates.
Box 6.2. Steps for estimating CDM\((m=0)\) using IPW estimation of MSMs with a categorical exposure and binary mediator

Step 1a. Obtain predicted exposure probabilities from a multinomial model for \(X\), unadjusted and adjusted for the exposure-outcome confounders, \(p(X = x)\) and \(p_{adj}(X = x)\), respectively.

Step 1b. Generate the weights corresponding to each level of the exposure

\[
w(X = x) = \frac{p(X = x)}{p_{adj}(X = x)}
\]

Step 2a. Obtain predicted mediator probabilities (unadjusted and adjusted for confounders),

\[
p(M = 1) = \{1 + \exp[-\alpha_0]\}^{-1}
\]

\[
p_{adj}(M = 1) = \left\{1 + \exp \left[ -\alpha_0 - \sum_{j=2}^{3} \alpha_{1j} I_{X=j} - \alpha'_2 C_{XY} - \alpha'_3 C_{MY} \right] \right\}^{-1}
\]

where \(I_{X=j}\) is an indicator of whether the exposure \(X\) takes value \(j\), \(C_{XY}\) are exposure-outcome confounders, \(C_{MY}\) are mediator-outcome confounders, bold indicates matrices and dashes ('\) indicate transposition of matrices.

Step 2b. Generate the weights for the mediator using the predicted values from these models, where the weights depend on the observed level of the mediator, e.g.

\[
w(M = 0) = \frac{p(M = 0)}{p_{adj}(M = 0)}
\]

Step 3. Generate overall weights by multiplying the exposure and mediator weights:

\[
ipw = w(X = x)w(M = m)
\]

Step 4. Fit a weighted regression model of the outcome against the exposure, the mediator and the interaction between the exposure and mediator, using the \(ipw\) weights:

\[
E_{ipw}(Y) = \theta_0 + \theta_1 X + \theta_2 M + \theta_3 XM
\]
Box 6.3. Steps for estimating $CDM(\text{m}=0)$ using $g$-estimation of structural nested models with a categorical exposure and binary mediator

Step 1a. Obtain predicted mediator probabilities from a logistic regression model for $M$, adjusted for the exposure-outcome and mediator-outcome confounders.

$$\hat{p}(M = 1) = \{1 + \exp\left[-\delta_0 - \sum_{j=2}^{3} \delta_{1j} I_{X=j} - \delta'_{2} C_{XY} - \delta'_{3} C_{MY}\right]\}^{-1}$$

Step 1b. Derive mediator residuals, $\hat{r}(M) = M - \hat{p}(M)$ where $\hat{p}(M)$ is the estimate of the propensity for $M = 1$.

Step 2. Regress the outcome against mediator residuals, the interaction between exposure and mediator residuals, and mediator-outcome confounders using ordinary least squares,

$$E(Y|X, M, C_{XY}, C_{MY}) = \beta_0 + \gamma_2 \hat{r}(M) + \sum_{j=2}^{3} \gamma_{3j} I_{X=j} \hat{r}(M) + \beta'_{1} C_{XY} + \beta'_{2} C_{MY}$$

Step 3. Create a transformed outcome by removing the estimated mediator effect from the observed outcome, $\tilde{Y} = Y - \hat{\gamma}_{2} M - \sum_{j=2}^{3} \hat{\gamma}_{3j} I_{X=j} M$

Step 4a. Obtain predicted exposure probabilities from a multinomial model for $X$, controlling for the exposure-outcome confounders, $C_{XY}$.

Step 4b. Create exposure residuals for the levels of the exposure that are not the reference (baseline) value, $\hat{r}_{2}(X) = I_{X=2} - \hat{p}(X == 2)$ and $\hat{r}_{3}(X) = I_{X=3} - \hat{p}(X == 3)$

Step 5. Regress transformed outcome against predicted exposure residuals and exposure-outcome confounders using ordinary least squares to estimate counterfactual disparity measure, $E(Y) = \theta_0 + \psi_{2} \hat{r}(X) + \hat{r}_{2}(X) + \hat{r}_{3}(X) + \theta_{2} M + \theta C_{XY}$
6.7. Results

6.7.1. Missing data

After exclusions were applied, there were 274,444 infants included in the birth cohort (Figure 6.4). Of these, 8,946 (3.3%) had missing information for at least one study variable. Table 6.4 shows the distribution of cohort characteristics by infants with and without missing data. Missingness was driven by maternity records, which were unavailable (i.e. unlinked) for 5,913 (2.2%) of infants. The highest proportion of missing information was for maternal smoking, for which 3.3% of infants did not have a record. Maternal age, maternal asthma and parity all had 2.2% of records missing. In the adjusted logistic regression model for the odds of record incompleteness, missing data on ≥1 variable was associated with non-northern areas of residence (particularly Glasgow), earlier year of birth, non-UK maternal country of birth and, to a lesser extent, lower SIMD groups (see Table 6.4). Using simulation studies, White and Carlin illustrate that in circumstances where missingness is independent of the outcome given the covariates, complete case analysis has negligible bias. Analysis in this study was therefore performed using the 96.7% of infants with complete data.

Figure 6.4. Flow diagram showing derivation of final study cohorts for mediation analysis and calculations of admission rates

*Child-IDs belonging to children born in 2017 or excluded from birth cohort at earlier stage of diagram
Table 6.4. Cohort characteristics, missing data and admission rates

<table>
<thead>
<tr>
<th></th>
<th>Infants</th>
<th>Complete cases</th>
<th>Odds of ≥1 missing value</th>
<th>Admissions per 1000 infant-years</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>All N %</td>
<td>N %</td>
<td>aOR** (95% CI)</td>
<td>Rate (95% CI)</td>
</tr>
<tr>
<td>Total</td>
<td>274,444 100.0</td>
<td>265,498 95.2</td>
<td>1.09 (0.98, 1.21)</td>
<td>55.5 (54.6, 56.4)</td>
</tr>
<tr>
<td>≥1 adm. for bronchiolitis</td>
<td>No</td>
<td>261,311 95.2</td>
<td>Reference</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Yes</td>
<td>13,133 4.8</td>
<td>Reference</td>
<td></td>
</tr>
<tr>
<td>Year of birth</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2012</td>
<td></td>
<td>56,835 20.7</td>
<td>1.67 (1.56, 1.79)</td>
<td>48.5 (46.7, 50.3)</td>
</tr>
<tr>
<td>2013</td>
<td></td>
<td>54,839 20.0</td>
<td>1.47 (1.37, 1.58)</td>
<td>47.9 (46.1, 49.8)</td>
</tr>
<tr>
<td>2014</td>
<td></td>
<td>55,414 20.2</td>
<td>1.08 (1.00, 1.16)</td>
<td>54.7 (52.8, 56.7)</td>
</tr>
<tr>
<td>2015</td>
<td></td>
<td>53,946 19.7</td>
<td>1.23 (1.15, 1.33)</td>
<td>58.5 (56.5, 60.6)</td>
</tr>
<tr>
<td>2016</td>
<td></td>
<td>53,410 19.5</td>
<td>Reference</td>
<td>68.8 (66.6, 71.0)</td>
</tr>
<tr>
<td>SIMD</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Low</td>
<td></td>
<td>67,585 24.6</td>
<td>1.05 (0.99, 1.11)</td>
<td>45.0 (43.4, 46.6)</td>
</tr>
<tr>
<td>Medium</td>
<td></td>
<td>106,268 38.7</td>
<td>1.13 (1.07, 1.18)</td>
<td>52.5 (51.1, 53.9)</td>
</tr>
<tr>
<td>High</td>
<td></td>
<td>100,591 36.7</td>
<td>Reference</td>
<td>65.7 (64.1, 67.3)</td>
</tr>
<tr>
<td>Area of residence</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Northern</td>
<td></td>
<td>34,884 12.7</td>
<td>Reference</td>
<td>41.4 (41.4, 45.8)</td>
</tr>
<tr>
<td>Central</td>
<td></td>
<td>78,119 28.5</td>
<td>1.19 (1.09, 1.30)</td>
<td>52.4 (50.8, 54.0)</td>
</tr>
<tr>
<td>Glasgow</td>
<td></td>
<td>64,671 23.5</td>
<td>3.18 (2.93, 3.46)</td>
<td>45.0 (43.4, 46.7)</td>
</tr>
<tr>
<td>Edinburgh</td>
<td></td>
<td>48,450 17.7</td>
<td>1.36 (1.24, 1.49)</td>
<td>58.6 (56.5, 60.8)</td>
</tr>
<tr>
<td>Southern</td>
<td></td>
<td>48,320 17.6</td>
<td>1.30 (1.18, 1.43)</td>
<td>79.9 (77.4, 82.5)</td>
</tr>
<tr>
<td>Maternal birth country</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Non-UK</td>
<td></td>
<td>43,728 15.9</td>
<td>1.35 (1.28, 1.43)</td>
<td>35.5 (33.7, 37.3)</td>
</tr>
<tr>
<td>UK</td>
<td></td>
<td>230,716 84.1</td>
<td>Reference</td>
<td>59.3 (58.3, 60.3)</td>
</tr>
<tr>
<td>Maternal smoking</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>No</td>
<td></td>
<td>215,704 81.2</td>
<td>Reference</td>
<td>49.7 (48.7, 50.6)</td>
</tr>
<tr>
<td>Yes</td>
<td></td>
<td>49,794 18.8</td>
<td>82.4 (80.0, 85.0)</td>
<td></td>
</tr>
<tr>
<td>Missing</td>
<td></td>
<td>8,946 3.3</td>
<td>Reference</td>
<td>47.7 (43.3, 52.5)</td>
</tr>
<tr>
<td>Maternal age (years)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>≤19</td>
<td></td>
<td>12,062 4.4</td>
<td>68.0 (63.5, 72.8)</td>
<td></td>
</tr>
<tr>
<td>20-29</td>
<td></td>
<td>119,111 43.4</td>
<td>72.6 (70.1, 75.2)</td>
<td></td>
</tr>
<tr>
<td>30-39</td>
<td></td>
<td>127,095 46.3</td>
<td>49.2 (47.7, 50.7)</td>
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</tr>
<tr>
<td>≥40</td>
<td></td>
<td>10,263 3.7</td>
<td>42.9 (39.8, 46.3)</td>
<td></td>
</tr>
<tr>
<td>Missing</td>
<td></td>
<td>5,913 2.2</td>
<td>Reference</td>
<td>48.8 (42.4, 54.8)</td>
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<tr>
<td>Maternal asthma</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>No</td>
<td></td>
<td>248,526 90.6</td>
<td>53.4 (52.5, 54.3)</td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td></td>
<td>20,005 7.3</td>
<td>84.0 (80.1, 88.1)</td>
<td></td>
</tr>
<tr>
<td>Missing</td>
<td></td>
<td>5,913 2.2</td>
<td>48.8 (43.4, 54.8)</td>
<td></td>
</tr>
<tr>
<td>Older siblings</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>No</td>
<td></td>
<td>113,783 41.5</td>
<td>36.4 (35.3, 37.5)</td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td></td>
<td>154,748 56.4</td>
<td>69.7 (68.4, 71.0)</td>
<td></td>
</tr>
<tr>
<td>Missing</td>
<td></td>
<td>5,913 2.2</td>
<td>48.8 (43.4, 54.8)</td>
<td></td>
</tr>
</tbody>
</table>

**(aOR=adjusted odds ratio) multivariable logistic regression comparing the odds of having at least one missing value compared with no missing values, adjusted for bronchiolitis admission, year of birth, socioeconomic deprivation, country of birth and area of residence**
6.7.2. Cohort description (complete cases)

There were 265,498 infants in the final cohort of which 12,766 (4.8%) had at least one hospital admission with a diagnosis of bronchiolitis in the first year of life. The proportion of children with at least one admission for bronchiolitis increased over time, from 4.3% for children born in 2012 to 5.8% for children born in 2016. Overall, 36.6% of infants were in the high SIMD group and 18.8% of mothers smoked during their pregnancy. When split by deprivation group, there are differences in the distribution of key risk factors across the cohort (Table 6.5). Notably, 29.2% of mothers in the high SIMD group reported smoking during pregnancy compared to 7.2% in the low SIMD group. 5.7% of infants in the high SIMD group were admitted to hospital with bronchiolitis at least once in the first year of life compared to 3.9% of infants in the low deprivation group. There were also marked differences by area of residence (30.3% of infants in the high SIMD group are resident in greater Glasgow at birth), maternal age (lower in the low SIMD group) and maternal asthma (highest in the high SIMD group).

6.7.3. Admission rates (all infants)

As shown in Table 6.4, the overall incidence rate of admissions for bronchiolitis over the study period was 55.5 per 1000 infant-years (95% CI 54.6 to 56.4). The rate of admissions increased over time, from 48.5 per 1000 infants-years (95% CI 46.7 to 50.3) in 2012 to 68.8 per 1000 infant-years (95% CI 66.6 to 71.0) in 2016, and was particularly high among children with a southern Scottish residence listed at birth (79.9 per 1000 infant-years, 95% CI 77.4 to 82.5). The rate of admissions increased with level of socioeconomic deprivation, with infants in the most deprived SIMD group having an admission rate of 65.7 per 1000 infant-years (95% CI 64.1 to 67.3) compared to 45.0 per 1000 infant-years (95% CI 43.4 to 46.6) in the least deprived SIMD group over the 5-year period. UK maternal country of birth, maternal smoking during pregnancy, younger maternal age, maternal asthma and having older siblings were all risk factors associated with higher rates of admissions.
### Table 6.5. Final cohort characteristics, by socioeconomic deprivation group, and ≥1 hospital admission for bronchiolitis in infancy

<table>
<thead>
<tr>
<th></th>
<th>SIMD group</th>
<th></th>
<th></th>
<th>≥1 bronchiolitis admission</th>
<th>N</th>
<th>% of row</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Low (N)</td>
<td>Medium (N)</td>
<td>High (N)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>≥1 bronchiolitis admission</td>
<td>No 62,932</td>
<td>98,200 95.5</td>
<td>91,645 94.3</td>
<td>-</td>
<td>142</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Yes 2,554</td>
<td>4,730 4.6</td>
<td>5,540 5.7</td>
<td>2,776 100.0</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Deprivation</td>
<td>Low 65,486</td>
<td>102,827 100.0</td>
<td>97,185 100.0</td>
<td>2,776 100.0</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Medium -</td>
<td>-</td>
<td>-</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>High -</td>
<td>-</td>
<td>-</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Year of birth</td>
<td>2012 13,097</td>
<td>21,080 20.5</td>
<td>20,214 20.8</td>
<td>2,325 4.3</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>2013 12,835</td>
<td>20,565 20.0</td>
<td>19,437 20.0</td>
<td>2,232 4.2</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>2014 13,359</td>
<td>20,771 20.2</td>
<td>19,729 20.3</td>
<td>2,585 4.8</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>2015 13,097</td>
<td>20,154 19.6</td>
<td>19,048 19.6</td>
<td>2,623 5.0</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>2016 13,097</td>
<td>20,227 19.7</td>
<td>18,757 19.3</td>
<td>3,001 5.8</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Maternal smoking</td>
<td>No 60,771 92.8</td>
<td>86,066 83.7</td>
<td>68,807 70.8</td>
<td>9,301 4.3</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Yes 4,715 7.2</td>
<td>16,761 16.3</td>
<td>28,378 29.2</td>
<td>3,465 7.0</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Area of residence</td>
<td>Northern 7,072 10.8</td>
<td>17,789 17.3</td>
<td>9,427 9.7</td>
<td>1,308 3.8</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Central 23,837 36.4</td>
<td>29,306 28.5</td>
<td>23,130 23.8</td>
<td>3,472 4.6</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Glasgow 13,097 20.0</td>
<td>18,303 17.8</td>
<td>29,447 30.3</td>
<td>2,442 4.0</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Edinburgh 15,389 23.5</td>
<td>18,509 18.0</td>
<td>13,314 13.7</td>
<td>2,306 4.9</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Southern 6,156 9.4</td>
<td>19,023 18.5</td>
<td>21,964 22.6</td>
<td>3,237 6.9</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Maternal age (years)</td>
<td>≤19 851 1.3</td>
<td>3,907 3.8</td>
<td>6,997 7.2</td>
<td>703 6.0</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>20-29 19,056 29.1</td>
<td>45,141 43.9</td>
<td>53,452 55.0</td>
<td>6,352 5.4</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>30-39 41,846 63.9</td>
<td>49,768 48.4</td>
<td>34,306 35.3</td>
<td>5,335 4.2</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>≥40 3,733 5.7</td>
<td>4,010 3.9</td>
<td>2,430 2.5</td>
<td>376 3.7</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Maternal asthma</td>
<td>No 61,360 93.7</td>
<td>95,321 92.7</td>
<td>89,021 91.6</td>
<td>11,400 4.6</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Yes 4,126 6.3</td>
<td>7,506 7.3</td>
<td>8,164 8.4</td>
<td>1,366 6.9</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Maternal birth country</td>
<td>Non-UK 10,281 15.7</td>
<td>15,321 14.9</td>
<td>16,327 16.8</td>
<td>1,295 3.1</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>UK 55,205 84.3</td>
<td>87,506 85.1</td>
<td>80,858 83.2</td>
<td>11,471 5.1</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Older siblings (parity)</td>
<td>No 27,308 41.7</td>
<td>44,524 43.3</td>
<td>39,749 40.9</td>
<td>3,609 3.2</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Yes 38,178 58.3</td>
<td>58,303 56.7</td>
<td>57,436 59.1</td>
<td>9,157 5.9</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
6.7.4. Mediation analyses

As shown in Table 6.6, after accounting for exposure-outcome confounders, the potential risk of at least one hospital admission for bronchiolitis in the first year of life rose with each level of deprivation (SIMD). The two approaches used to calculate potential risk and risk differences produced almost identical results and I present the IPW of MSMs-derived estimates here. After accounting for year of birth, maternal country of birth and area of residence by IPW estimation, the potential risk of bronchiolitis admission in infancy would be 40.8 per 1000 live births (95% CI 39.2 to 42.3) if everyone were assigned to the low SIMD group at birth. This compares to 57.1 per 1000 live births (95% CI 55.6 to 58.6) if all infants were assigned the high SIMD group. Overall for every 1000 live births, if all children were assigned to the high SIMD group, there would be 16.3 additional infants with (at least one) admission for bronchiolitis per 1000 live births (95% CI 14.4 to 18.2). The additional potential risk would be 4.6 per 1000 infants were all infants assigned to the medium SIMD group (95% CI 2.4 to 6.8).

IPW of MSMs suggested that the counterfactual disparity between the high and low SIMD groups would be reduced to 13.0 per 1000 live births (95% CI 10.0 to 16.0) if maternal smoking were eliminated. This translates into a 20.3% disparity reduction in admissions (95% CI 12.1 to 28.4). A slightly lower reduction in counterfactual disparity was estimated using g-estimation of structural nested models (13.4 potential risk difference, 95% CI 11.0 to 15.7; 18.0% disparity reduction, 95% CI 10.3 to 25.6). The estimated remaining disparity in the medium compared to the low SIMD group was 3.3 per 1000 live births (95% CI 1.2 to 5.5) using the IPW of MSMs method. This would be a disparity reduction of 27.2% (95% CI 9.0 to 45.4) if maternal smoking were eliminated. The g-estimation of MSMs method suggested a remaining disparity of 4.0 admissions per 1000 live births (95% CI 1.8 to 6.1), which translated to noticeably smaller disparity reduction of 14.5% (95% CI -9.1 to 38.2) between the medium and low SIMD groups.
Table 6.6. Potential risk of ≥1 admission to hospital for bronchiolitis during infancy, estimated counterfactual disparity in admissions and corresponding disparity reduction explained by maternal smoking, by g-method and SIMD group

<table>
<thead>
<tr>
<th>SIMD group</th>
<th>Marginal risk per 1000 live births* (95% CI)</th>
<th>Marginal risk difference per 1000 live births* (95% CI)</th>
<th>CDM(m=0) per 1000 live births (95% CI)**</th>
<th>% disparity reduction (95% CI)**†</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>IPW MSMs</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Low</td>
<td>40.8 (39.2, 42.3)</td>
<td>Reference</td>
<td>Reference</td>
<td>Reference</td>
</tr>
<tr>
<td>Medium</td>
<td>45.4 (44.1, 46.7)</td>
<td>4.6 (2.4, 6.8)</td>
<td>3.3 (1.2, 5.5)</td>
<td>27.2 (9.0, 45.4)</td>
</tr>
<tr>
<td>High</td>
<td>57.1 (55.6, 58.6)</td>
<td>16.3 (14.4, 18.2)</td>
<td>13.0 (10.0, 16.0)</td>
<td>20.3 (12.1, 28.4)</td>
</tr>
<tr>
<td><strong>G-estimation of SNMs</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Low</td>
<td>40.7 (39.1, 42.3)</td>
<td>Baseline</td>
<td>Baseline</td>
<td>Baseline</td>
</tr>
<tr>
<td>Medium</td>
<td>45.4 (44.2, 46.5)</td>
<td>4.6 (2.5, 6.8)</td>
<td>4.0 (1.8, 6.1)</td>
<td>14.5 (-9.12, 38.2)‡</td>
</tr>
<tr>
<td>High</td>
<td>57.0 (55.2, 58.9)</td>
<td>16.3 (14.1, 18.5)</td>
<td>13.4 (11.0, 15.7)</td>
<td>18.0 (10.3, 25.6)</td>
</tr>
</tbody>
</table>

*Risk and absolute risk difference after adjusting for differences in year of birth, maternal country of birth and area of residence; **95% CIs are calculated using the bootstrap with 1000 replications; †disparity reduction (%) was estimated as = (risk difference-CDM(m=0))/risk difference*100; ‡bootstraps of the g-estimation procedure produced some estimates of the CDM that were larger than the TCE, leading to some negative percentages.

6.8. Discussion

Using a population-based Scottish birth cohort, I employed a counterfactual causal inference approach to establish the extent to which maternal smoking explains the socioeconomic disparity in the risk hospital admission for bronchiolitis in infancy. The overall relationship between SIMD and bronchiolitis admission was 16.3 (95% CI 14.4 to 18.2) more children with at least one hospital admission for bronchiolitis in infancy for every 1000 infants born into high compared to low levels of SIMD. Using the CDM approach, it was estimated that the elimination of maternal smoking would reduce about one fifth of the disparities in the proportion of children who had at least one hospital admission for bronchiolitis in the first year of life.

6.8.1. Strengths and weaknesses

Linked Scottish health datasets

The coverage and completeness of the birth cohort used in this study is a major strength of this work. The core cohort was derived from vital records held by NRS, meaning that almost all infants born in Scotland between 2012 and 2016 were included in the study. Infant information was supplemented with maternal delivery records, dispensing information and hospital admission data, creating a comprehensive dataset. This enabled me to capture
important risk factors in the development of offspring respiratory illness that are not available in the equivalent English dataset, including indicators of maternal asthma and maternal smoking during pregnancy. However, there are also drawbacks to this dataset.

Whilst variables recorded in NRS birth registrations were 100% complete, 2.2% of children’s records were not linked to maternal records, meaning that they was missing information on several key variables. Maternal ethnicity was particularly affected by missingness and was therefore not included in the main analysis despite being identified as a potential confounder between maternal smoking and offspring bronchiolitis. Omission of ethnicity in Scottish hospital records is a long noted problem, although recording of this characteristic has improved markedly since 2010 when the Equality Act was brought in (from 42% completeness in 2010 to 82% in 2016). Other studies using Scottish administrative data have benefitted from linkage to census data for higher completeness of ethnicity.

I did include a related variable (maternal country of birth) as a confounder, which likely partially captured the confounding effect of maternal ethnicity. Although I used a structured method to derive study confounders, there may also be additional unmeasured confounding, which would violate the conditional exchangeability assumption.

Measurement error is likely present in the study mediator. Maternal smoking is self-reported in this study and, as a factor known to be affected by social desirability bias, is likely underreported. A study of pregnant women in 2003/04 in the West of Scotland found that self-reported smoking status from SMR02 (as used in this study) underestimated smoking by 25% when compared to an objective measurement of smoking, and women from more affluent areas were more likely to misreport their smoking status. In addition, having a singular binary indicator of maternal smoking measured at one time point (during pregnancy) means that the full complexities of this risk factor, including the amount and timing of tobacco smoke exposure, could not be studied. In particular, many women who smoke during pregnancy are likely to continue after birth, so it is difficult to differentiate between the effects during pregnancy and smoking exposure after birth. Conversely, there will be some non-smokers who live with someone who smokes in the house. Linkage to datasets that capture early exposure to second hand smoke, such as the Child Health Surveillance Programme Pre-School, would allow for more nuanced analyses of the effect of smoking on early respiratory health.

The PIS database does not include diagnosis codes and very few mothers had hospital admissions with a diagnosis of asthma, meaning that the indicator of maternal asthma used in this study relied primarily on dispensed medication data, which could have resulted in disease misclassification. Other studies have identified chronic obstructive pulmonary disease
as a source of misclassification with asthma; however as this condition is relatively uncommon in people younger than 40, this is unlikely to be a large source of bias in this study. Given that the pathway between maternal asthma and child bronchiolitis is far from determined—and likely includes a mixture of a genetic predisposition to respiratory illness and shared environmental factors—the use of an indicator that may have captured a broader group of maternal respiratory ill-health (as in this study) is unlikely to be of detriment to the findings.

**ESC-DAG (Objective 6B)**

The ESC-DAG protocol helped me to follow a systematic method to search the literature and create a study DAG. This evidence-based DAG construction and its application to DAGitty (which identifies causal and biasing paths) gives me confidence in the adjustment strategy for my mediation analysis. The ESC-DAG process gave me the opportunity to re-examine how previous research describing inequities in bronchiolitis have considered covariates in their analyses. Commonly, I reversed the directed edges between pathways that were treated as confounders in the original research but were likely mediators. This highlights a common issue of over-control for mediators in these studies, which may have led to an underestimation of the impact of socioeconomic deprivation on bronchiolitis rates.

There were downsides to using an ESC-DAG approach. Prominently, it was a particularly time intensive undertaking. Despite a restrictive exclusion criteria for the literature search, which resulted in the inclusion of 9 studies, I still had to undertake 552 directed edges assessments. I used a formal method to assess directed edges, considering temporality, face-validity, prior theory/research and counterfactual outcomes, but this did not completely eliminate subjectivity in the process. Consider, for example, the pathway from area of residence to number of siblings. Family size does vary by area in the UK, but it is unclear whether there is any direct link between the two factors or whether this association could be completely explained by alternative correlated factors such as socioeconomic position and ethnicity. For the most part, I erred on the side of caution for these debatable cases; keeping the directed edge rather than removing it (which would be a stronger assumption). The reliance on previous research to guide variable selection meant that understudied research areas, particularly paternal risk factors, were not included in the ESC-DAG.

Future applications may benefit from independent assessment of decisions using a secondary reviewer. In addition, a simplified version of the ESC-DAG process — for example, by combining related covariates into one node for assessment or excluding covariates unrelated to the mediator at an earlier stage in the process — is advisable where time is constrained. The
process took many days and may only be feasible for large projects and/or scenarios where there is a very simple and specific research question.

*Causal inference approach (Objectives 6A & 6C)*

I outlined the main mediation estimand(s) available and chose CDM as the most appropriate to answer my research question, given the available data and complexities of estimation. This estimand enabled me to characterise the overall level of disparity and capture the proportion of the disparity in the percentage of children with at least one bronchiolitis admission due to the impact of socioeconomic deprivation that would remain if maternal smoking were intervened upon. A major advantage of this approach is that, in comparison to other estimands, fewer assumptions were required to obtain unbiased estimates of the CDM. Several assumptions were still required, however, including no unmeasured mediator-outcome confounding, no interference, positivity and conditional exchangeability. I clarified how most of these assumptions were met in this study in section 6.2.2. The small number of confounders and broad categories used to define them (e.g. UK/non-UK for maternal country of birth) gives me confidence that the positivity assumption was met.

However, there are also some drawbacks of the CDM approach. In order to apply this approach I needed to alter my original research question to specify that the disparity measure was based on the scenario of no women smoking during pregnancy. This assumes an intervention that leads to a 100% stopping smoking rate that, based on the outcome of current interventions, is arguably unrealistic. One stop smoking service for pregnant women in a deprived area of Scotland, for example, achieved a 32% stop smoking rate at 4 weeks after the participants’ quit dates.425 Other estimands, such as the RIA-NIE that allows the mediator to be set to take a value that is randomly drawn from a distribution, arguably offer a more policy relevant estimate. However, this method could not be applied with the assurance of unbiased estimates.

I used two approaches to estimate the magnitude of socioeconomic disparity in infant hospital admission for bronchiolitis that would remain if no women smoked during pregnancy. The similarity in results for the high compared to low groups across the two methods gives me assurance of the robustness of these estimations. Both confidence intervals for these estimates are wide, which may be expected when the calculation involves the ratio of two risk differences; however, a percentage disparity estimate that is lower than 0% (as was the lower bound of the g-estimation of structural nested models method) is less plausible. It has been noted elsewhere that MSMs cope well with difference outcome variables, including logistic
regression (as was applied in this study), whereas there are issues applying structural nested models to binary data.426

6.8.2. Research findings and implications (Objective 6D)

Overall, this work shows that hospital admission rates for bronchiolitis have continually increased in Scotland. From 2012 to 2016, the rate of admissions increased from 48.5 per 1000 infant-years to 68.8 per 1000 infant-years. A similar difference is apparent between socioeconomic groups; infants from the least deprived socioeconomic group had an admission rate of 45.0 per 1000 infant-years compared to 65.7 per 1000 infant-years in the most deprived group across the whole study period. These findings continue to highlight the importance of work to tackle this condition and unravel the social determinants of inequity in admission rates.15

Looking beyond descriptive statistics, the main purpose of this study was to apply formal mediation methods to look at the specific pathway of maternal smoking during pregnancy in the association between socioeconomic position and hospital admission for bronchiolitis. Previous studies in this field have looked at the adjusted association of several risk factors on early-life RSV-related illnesses without focussing on elucidating socioeconomic pathways.44,137,138 I estimate that about one fifth of the difference in being admitted to hospital for bronchiolitis between infants in high and low socioeconomic groups in Scotland can be attributed to maternal smoking. My results therefore show the continuing need to reduce rates of maternal smoking specifically in the most socioeconomically deprived populations. These reductions would substantially reduce the social gradient in bronchiolitis hospital admissions.

With the rapid advances in methodology in the field of causal mediation analysis, future studies will likely benefit from methods that can better account for follow-up time and multiple admissions by the same child. In addition, future studies using similar formal methods but focussing on other policy-relevant mediators, such as outdoor air pollution and indoor conditions, such as damp and mould, are extremely important to deciphering the development of bronchiolitis and positioning interventions with the greatest impact.

6.9. Chapter summary

In this chapter, I applied a counterfactual causal inference approach to quantify the role of maternal smoking in bronchiolitis hospital admission disparities. To achieve this I have reviewed a broad range of mediation estimands recently proposed in the literature and discussed the potential application of these to my research question. I also employed a novel
and evidence-based approach to inform the appropriate adjustment strategy for my study to minimise bias. I included maternal country of birth, year, maternal age, maternal asthma, parity and area of residence as confounders of the mediator-outcome relationship. I found that an intervention aimed at eliminating maternal smoking during pregnancy could reduce the disparities in hospital admissions for bronchiolitis between the most and least deprived infants by up to one fifth.
Chapter 7. Trajectories of asthma/wheeze symptoms and their association with early-life risk factors

Recap and chapter overview

Chapters 4 to 6 explored socioeconomic inequities in bronchiolitis admissions among infants in England and Scotland. I used harmonic regression analyses to investigate the association between the seasonality of bronchiolitis admissions and socioeconomic deprivation at both the area and individual level. I then applied a counterfactual causal inference approach to quantify the role of maternal smoking in bronchiolitis hospital admission disparities.

In this chapter, I move my focus to childhood asthma; a common and chronic respiratory condition that has been linked to early-life bronchiolitis infection. Applying latent growth modelling techniques to the Scottish health datasets, I identify trajectories of asthma/wheeze symptoms among children aged less than 10 years. The association between early risk factors and asthma/wheeze group membership are then explored. Lastly, drawing from the counterfactual disparity work presented in Chapter 6, I investigate the mediating contribution of bronchiolitis in the pathway between socioeconomic position and asthma/wheeze. The final two thesis objectives are addressed in this chapter:

- To describe typical trajectories of asthma/wheeze among children <10 years in Scotland and their association with early life risk factors, including socioeconomic position
- To investigate the mediating contribution of bronchiolitis in the pathway between early life SEP and mid-childhood asthma/wheeze

7.1. Background

In 2018, 8% of children younger than 15 years old and 17% of those aged 16 to 24 years in Scotland reported a current doctor diagnosis of asthma. Despite this high prevalence, substantial contribution to morbidity and associated health-care burden across the world, the precise causes of asthma are relatively unknown. Moreover, the last decade has seen a stagnation in improvements of clinical outcomes, including mortality rates, for people with the condition. Asthma experts from a multitude of medical disciplines, writing in a 2018 Lancet
comission, argue that the lack of distinction between the different disease phenotypes under the umbrella term of “asthma” is a major contributor to this stagnation. These experts placed strong emphasis on the need to move beyond the current over-simplistic definition of asthma, which is currently based on common physiology and symptoms, despite encompassing conditions with pathobiologically distinct mechanisms.

Cohort studies in the UK, USA, the Netherlands and Taiwan, as well as in other countries, have been used to investigate different patterns of symptoms (often referred to as phenotypes) that comprise childhood asthma. Using statistical models that allow researchers to identify typical sub-groups of symptoms in longitudinal cohort data, between 3 and 6 trajectory groups have been identified based on the prevalence of asthma and wheeze symptoms at different ages. When defined by symptom onset and longevity, these groups are broadly characterised as:

- never/infrequent asthma and/or wheeze;
- early-transient asthma/wheeze, beginning before age 3 and subsiding after a few years;
- early-persistent asthma/wheeze, beginning before 3 and persisting into early adolescence (or the length of study follow-up);
- intermediate-onset asthma/wheeze, beginning between 3 and 5 years and persisting beyond study follow-up;
- and late-onset asthma/wheeze, beginning between 6 and 8 years and persisting beyond study follow-up.

The precise number of groups identified varies between studies, depending on factors such as the length of follow-up, definitions of asthma/wheeze employed and the sample size. As these groups are not directly observable but, rather, constructed from the data, they are described as latent variables. However, this does not mean that these different groupings are not recognised clinically. This data driven approach to classification has found to be comparable with clinical phenotypes defined by patient histories, diagnostic work-up and treatment responses.

Once trajectory groups have been identified, the probability of a study participant following each trajectory can then be associated with risk factors commonly associated with childhood respiratory ill-health, such as prenatal smoking exposure and early respiratory infections. Differentiating between asthma/wheeze groups and understanding trajectory-specific risk factors can better inform prognosis, and may be critical to identifying for which children and at which time early interventions may be best place to halt the progression of this disease into adulthood. Further, such analyses may be key to untangling the complex relationship
between socioeconomic deprivation and asthma, particularly the pathway relating bronchiolitis to asthma. As discussed in Chapter 1, this relationship is complex because of the likely interplay of environmental and biological risk factors occurring prenatally, in-utero and in early life on the development of an infant’s respiratory functions.47

Limitations have been noted in asthma trajectory studies to date, however. A 2020 systematic review and meta-analysis of childhood wheeze trajectory-specific risk factors identified high risk of information bias in all of the 13 included cohort studies.197 Child or parental reports, rather than clinical validation of symptoms, limited sample sizes and loss to follow-up are noted problems that lead to low power to detect associations with risk factors.431 Overcoming many of these previous shortcomings, Sbihi et al.24 used administrative health data to identify different asthma trajectories in a sample of more than 65,000 children in British Columbia. Results of Sbihi et al.’s study demonstrate the unique opportunity administrative data offers to study asthma trajectories, overcoming logistic and financial constraints and self-report biases inherent to purposefully designed longitudinal cohort studies based on questionnaires or interviews.24

The aim of this Chapter is to identify trajectories of asthma/wheeze symptoms in a Scottish context and to study whether and how they are socially patterned.

The objectives for this chapter are threefold:

**Objective 7A** To identify typical trajectories of asthma/wheeze in children aged <10 years old using Scottish administrative health data.

**Objective 7B** To examine how the identified asthma/wheeze trajectory groups vary according to social and biological risk factors operating early in the life course.

**Objective 7C** To quantify the mediating role of bronchiolitis in the pathway between socioeconomic position and the asthma/wheeze trajectory groups.

## 7.2. Study cohort and variable selection

### 7.2.1. Study DAG for mediation analyses

To identify confounders of the mediator-outcome (bronchiolitis admissions-asthma/wheeze) relationship, I adapted the diagram that emerged from the ESC-DAG procedure in Chapter 6 (Figure 6.1). Using the research presented in my introductory chapter, I determined that all identified risk factors for bronchiolitis were also associated with asthma and therefore, using
DAGitty, I drew a line from each node to asthma. To this, I added other risk factors for both bronchiolitis and asthma identified through my literature review in Chapter 1: passive smoking, and mould and damp. The assumed DAG for the mediator-outcome relationship is presented in Figure 7.1. This presents the minimal adjustment set to satisfy the assumption of no unmeasured mediator-outcome confounding: air pollution, area of residence, birth season, birth weight, birth year, breastfed, breech presentation birth, bronchopulmonary dysplasia, congenital anomalies, delivery method, child ethnicity, gestational age, gestational diabetes, maternal age, maternal asthma, maternal country of birth, maternal smoking, parity, preeclampsia, child sex and small for gestational age (SGA). As shown using pink nodes, some of these confounders are unavailable in my dataset. The implications of this are reviewed further in the discussion.
Figure 7.1. Assumed causal DAG for the mediator-outcome (bronchiolitis admission-chronic asthma) association.

Confounders that are available in my dataset are indicated by white nodes and unavailable confounders by red nodes.

The yellow node is the mediator and blue node is the outcome.
7.2.2. Data sources and study cohort

I used a subsection of the birth cohort of Scottish children described in Chapter 3 as the cohort for this study. The linked datasets that formed the cohort as well as covariates relevant to this study are shown in Table 7.1. In brief, information on births and deaths were obtained from NRS, with the birth register forming the core spine of the cohort. The birth and death registration data were supplemented with information from SBR, a singular electronic record capturing all the neonatal care received by babies in Scotland, and information on emigrations from the CHI registry. Further information about the birth was contained in the mother’s delivery record, which was retrieved from SMR02—a dataset submitted by maternity hospitals on inpatient and day case activity relating to obstetrics. Subsequent hospital admissions among children in the cohort were retrieved from SMR01, which includes information on all inpatients and day cases discharged from hospitals in Scotland. Data were also extracted from the PIS, which contains information on medicines prescribed and dispensed within the community setting in Scotland.

Table 7.1. Scottish datasets used to create the study cohort in this study

<table>
<thead>
<tr>
<th>Acronym</th>
<th>Dataset</th>
<th>Information</th>
<th>Key variables</th>
</tr>
</thead>
<tbody>
<tr>
<td>NRS</td>
<td>National records of Scotland birth &amp; death registrations</td>
<td>Child</td>
<td>Postal area, SIMD, date of birth, date of death, child sex, date of death, country of birth (mother)</td>
</tr>
<tr>
<td>CHI registry</td>
<td>Community Health Index registry</td>
<td>Child</td>
<td>Date of emigration</td>
</tr>
<tr>
<td>SBR</td>
<td>Scottish birth register</td>
<td>Child</td>
<td>SIMD, birthweight, diagnoses</td>
</tr>
<tr>
<td>SMR02</td>
<td>Scottish morbidity records – obstetric related hospital admissions</td>
<td>Mother</td>
<td>Smoking status, age at delivery, parity, SIMD, method of delivery, birthweight, gestational age, maternal age</td>
</tr>
<tr>
<td>SMR01</td>
<td>Scottish morbidity records–general hospital admissions</td>
<td>Child and mother</td>
<td>Diagnoses, data of admission, date of discharge</td>
</tr>
<tr>
<td>PIS</td>
<td>Scottish national prescribing information system</td>
<td>Child</td>
<td>Medication name, date dispensed</td>
</tr>
</tbody>
</table>

The initial study population was all live births between 1st January 2007 and 31st June 2008 in Scotland. Follow-up began at birth and continued until the child’s 10th birthday. I excluded children who died or emigrated out of Scotland by 2 years of age. This step was taken to ensure that all children in the cohort had a recording of the study outcome (i.e. presence or no presence of asthma/wheeze symptoms) for at least one time point. Children born to
mothers of non-Scottish residency (identified through mother’s country of residency in the NRS birth registrations) were also excluded from the dataset to prevent potential systematic loss to follow up. One child from each non-singleton birth was randomly selected to be included in the study.

7.2.3. Study variables

Definition of asthma/wheeze

The outcomes in this study are trajectory groups of asthma/wheeze, which are determined through the latent growth curve modelling described in section 7.3.2, below. In order to model trajectories, I firstly needed to define instances of asthma/wheeze at specific time points. I used a multidimensional definition, with expert input from Steve Cunningham, Professor of Paediatric Respiratory Medicine at the University of Edinburgh, to ensure this classification reflected medical practice. For each year of age between 2 and 9, inclusive, children were defined as having asthma and/or wheeze if they met any of the following criteria in the 12-month period between birthdays:

- ≥1 hospital admission in SMR01 with a main diagnosis, as defined by International Classification of Diseases, 10th revision (ICD-10) codes, for asthma (J45 including all subcategories), status asthmaticus (J46X) or wheezing (R06.2);
- or ≥4 dispensed prescriptions for any specified asthma medication (see Table 7.2 for medications and relevant age exclusions);
- or asthma mentioned in death records as primary or secondary reason of death.

Asthma medication was defined using the British National Formulary for Children (BNFC),\textsuperscript{432} with age of child at each asthma/wheeze event was retrieved from the relevant dataset (SMR01, PIS or death record). I selected age 2, rather than an earlier age, as the start of symptom follow-up to reduce the risk of capturing instances of bronchiolitis (the study mediator) in the outcome.

Diagnosis of asthma is not recommended in children younger than 5, in part due to the difficulty of using standardised tests in this population.\textsuperscript{3} Moreover, recurrent symptoms of wheeze (typically defined as a high pitched sound with a musical quality that probably results from turbulence through narrowed tubes)\textsuperscript{433} is common in young children and is not necessarily an indicator of later asthma.\textsuperscript{434} As such, including symptoms between ages 2 and 4 in this study is intended to capture early wheeze patterns rather than asthma per se. Conversely, a child may have asthma, but currently be asymptomatic (thus not presenting with symptoms of
wheeze). It is for these reasons that I use the broader term “asthma/wheeze” rather than singular terms of asthma or wheeze to describe trajectories of symptoms in this study.

Table 7.2. Drugs that are used to treat asthma, defined using the BNFC

<table>
<thead>
<tr>
<th>BNF Chapter section</th>
<th>Conditions treated in children</th>
<th>Inclusions (age restriction applied)</th>
<th>Exclusions (age restriction applied)</th>
</tr>
</thead>
<tbody>
<tr>
<td>3.1 Bronchodilators</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>3.1.1.1 Selective beta&lt;sub&gt;2&lt;/sub&gt; agonists</td>
<td>Short acting beta2 agonists: Reversible airway obstruction Long acting beta2 agonists: Croup – single dose of corticosteroids</td>
<td>Salbutamol, Terbutaline Formoterol (age 6+), Salmeterol</td>
<td></td>
</tr>
<tr>
<td>3.1.1.2 Other adrenoceptor agonists</td>
<td>Acute allergic and anaphylactic reactions, angioedema, cardiopulmonary resuscitation, severe croup</td>
<td>Orciprenaline sulfate (withdrawn from the market in 2010)</td>
<td>Adrenaline (epinephrine), Ephedrine (hydrochloride)</td>
</tr>
<tr>
<td>3.1.2 Antimuscarinic bronchodilators</td>
<td>Reversible airway obstruction, rhinitis</td>
<td>Ipratropium bromide</td>
<td>Aclidinium bromide, Glycopyrronium, Indacaterol, Tiotropium, Umeclidinium</td>
</tr>
<tr>
<td>3.1.3 Theophylline</td>
<td>Reversible airway obstruction, bronchospasm associated with chronic bronchitis, neonatal apnoea</td>
<td>Theophylline, Aminophylline</td>
<td></td>
</tr>
<tr>
<td>3.1.5 Peak flow meters, inhaler devices and nebulisers</td>
<td></td>
<td>Standard range peak flow meter (age 6+), Low range peak flow meter (age 6+), Inhaler devices, Spacer devices</td>
<td></td>
</tr>
<tr>
<td>3.2 Corticosteroids</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>3.2 Corticosteroids</td>
<td>Croup, bronchopulmonary dysplasia</td>
<td>Beclometasone dipropionate (age 2+), Budesonide (age 6 +), Ciclesonide (age 12+), Fluticasone propionate (age 4+), Mometasone furoate (age 12+)</td>
<td></td>
</tr>
<tr>
<td>3.3 Cromoglicate and related therapy and leukotriene receptor antagonists</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>3.3.1 Cromoglicate and related therapy</td>
<td>Food allergy, allergic conjunctivitis, allergic rhinitis</td>
<td>Sodium cromoglicate (age 5+), Nedocromil Sodium (age 5+)</td>
<td></td>
</tr>
<tr>
<td>3.3.2 Leukotriene receptor antagonists</td>
<td>Allergic rhinitis (15-19 years)</td>
<td>Montelukast</td>
<td>Zafirlukast (12+ years only)</td>
</tr>
<tr>
<td>3.4.2 Allergen Immunotherapy</td>
<td></td>
<td>Omalizumab (6+)</td>
<td></td>
</tr>
</tbody>
</table>
6.3.2 Glucocorticoid therapy

| Glucocorticoid therapy | Croup, inflammation, ulcerative colitis, Crohn's disease | Hydrocortisone, Dexamethasone, Prednisone, Prednisolone (in conjunction with hospital admission) | Betamethasone, Deflazacort, Methylprednisolone |

**Exposure**

I used Scottish Index for Multiple Deprivation (SIMD) 2006 to capture socioeconomic position for each cohort member. SIMD 2006 deciles were split into three groups for analyses indicating high (top 30% rank of SIMD scores), medium (middle 40% rank of SIMD scores) and low levels of socioeconomic deprivation (bottom 30% rank of SIMD scores). This is the same exposure as used in Chapter 6, except SIMD 2006 deciles rather than SIMD 2012 deciles were used (because these are nearest indices to the birth years of cohort members). Again, SIMD deciles were retrieved from NRS birth registration files and supplemented from SBR or SMR02 where missing.

**Mediator**

I identified children with ≥1 hospital admission with a primary or secondary diagnosis of bronchiolitis during the first year of life in SMR01 records using ICD-10 code J21 for acute bronchiolitis. This was the same definition of bronchiolitis as used in Chapter 6.

**Confounders**

The selection of variables in this section are based on the confounders identified in the study DAG for mediation analysis (Figure 7.2, below). I refer to these as risk factors in the multivariable logistic regression models and as confounders in the mediation analysis.

Child sex (female/male), date of birth, mother’s country of birth (dichotomised into UK or non-UK born) and postcode area were retrieved from the child’s birth registration file. Year and month of birth, used as an indicator of birth season, were derived from date of birth. The 16 postcode areas in Scotland were grouped to make 5 broader areas with similar population sizes: northern Scotland, central Scotland, Glasgow, Edinburgh and southern Scotland (see Figure 6.3). I created an indicator of chronic conditions, including congenital anomalies and bronchopulmonary dysplasia, using the list of ICD-10 codes previously defined by Kristensen and colleagues (see Table A7.1, Appendix 7.1). Children were categorised as having a chronic condition (yes or no) if they had any of these diagnoses recorded in their birth record,
death record or any hospital admission before 6 months of age. The mother’s delivery record
provided method of delivery (categorised into vaginal, elective caesarean and emergency
caesarean), maternal age, parity, mother’s smoking status during pregnancy, gestational age
in weeks and birthweight in grams. Birthweight was supplemented with information from the
child’s birth record where missing.

Maternal age was coded into a four-category variable (< 20, 20–29, 30–39 and ≥40 years) and
parity (count of previous births) was used to create a number of older siblings variable that
was split into groups (0, 1 and 2 or more siblings). Mothers self-reported smoking status,
recorded at around 8-12 weeks gestation, was treated as a binary variable with yes signifying
mothers smoked traditional cigarettes (i.e. non-e-cigarettes) during their pregnancy.
Gestational age was categorised as preterm (<37 weeks) or non-preterm (≥37 weeks). SGA
(yes/no) was defined by birthweights falling below the 10th centile of the sex-, and gestational
age-specific distributions (using LMS tables).436 I created an indicator of severe maternal
asthma (yes/no), for mothers who had one or more hospital admission before their child’s birth
that included a diagnosis of asthma (ICD-10 code J45, including all subcategories).

Additional covariates

These additional variables were extracted from the datasets to be included as predictors in
the imputation models for missing data (see motivation for this in Sections 7.3.3 and 7.4.1).
The 9-class National Statistics Socio-Economic Classification (NS-SEC), based on father’s
occupation for married/cohabiting couples and on mother’s occupation for all other birth
registration types,437 was retrieved from NRS birth registration files. A binary intensive care
stay variable was generated from information in the child’s SBR file, with yes indicating any
postnatal intensive or high dependency care. Any child admitted to hospital diagnosed with
any of the following ICD-10 codes in the first few months of life were recorded as having a
prematurity-related complication: H35.1 (retinopathy of prematurity); P01.1 (newborn affected
by premature rupture of membranes); P07.0 (extremely low BW <1000g) and P07.2 (extreme
prematurity, <28 w); P27.0, P27.1, P27.8 and P27.9 (chronic respiratory diseases originating
in the perinatal period); P59.0 (neonatal jaundice associated with preterm delivery); and P61.2
(anemia of prematurity). This indicator was used in addition to the chronic conditions indicator
discussed above specifically for imputation purposes. Birth hospital was retrieved from the
child’s birth file and supplemented with the mother’s delivery file where missing.
7.3. **Statistical methods**

7.3.1. Overview of the analytical steps to address the 3 objectives

During the process of the four analytical steps in this chapter, I switch between statistical software. To illustrate the order in which these steps are carried out I have included a diagram of the process in Figure 7.2, below. Step 1 involved modelling asthma/wheeze trajectories (addressing objective 7A). In step 2, I used multiple imputation by chained equations (MICE) to impute missing values in risk factors (a process conducted independently of step 1). The results of step 1 and 2 were then combined to carry out multinomial logistic regression in step 3 (to examine characteristics of asthma/wheeze trajectory groups, objective 7B). For the mediation analysis (step 4), the classes derived from the final latent growth model were merged with the original (unimputed) dataset, then imputation, bootstrap inference and mediation analysis carried out together. I used Stata 15\textsuperscript{314} for descriptive data analysis, multiple imputation and mediation analysis and Mplus version 8.2\textsuperscript{438} to implement latent growth modelling and univariable multinomial logistic regression.

![Figure 7.2. Analysis steps in this Chapter, split by software](image-url)
7.3.2. Typical trajectories of asthma/wheeze (objective 7A)

To model asthma/wheeze trajectories, I used two approaches latent growth curve modelling: latent class growth analysis (LCGA), which is also known as group-based trajectory modelling, and growth mixture modelling (GMM). Both methods offer statistical approaches to detecting classes of individuals with a similar developmental trajectory, but differ in whether within-class heterogeneity is allowed. LCGA assumes there is no variation within a class, meaning that all individual trajectories within a class are represented by a single curve. GMM, on the other hand, allows variation across individuals within classes and produces a class of trajectories with a common mean and shape. As described by Muthén, GMM therefore offers “a more realistic representation of complex data” and, unlike with LCGA, the resultant classes may (cautiously) be interpreted as underlying phenotypes. However, GMM is also far more computationally intensive and noted for having convergence issues because of maximum likelihood estimation problems.

The purpose of latent growth curve modelling in this study was to model the probability of asthma/wheeze symptoms over time, allowing for different trajectories for these probabilities corresponding to the latent classes. I fitted both LCGA and GMMs. In each case, probabilities were modelled using a mixture of logistic distributions for the dichotomous outcomes of asthma/wheeze presence (yes/no) at the ages from 2 to 9 years, independently of other covariates. For GMMs, the logistic models have additional terms capturing the within class variability. The outcome, asthma/wheeze, was set to missing for all follow-up years following the year of emigration or death (for children with these emigration or death recorded before age 9). Mplus assumes that this missing data in the outcome is missing at random and uses information held in the repeated outcome observations to predict the missing values. Model selection requires comparing goodness-of-fit statistics obtained from models fitted with varying numbers of trajectory groups and shapes. I began my analyses with LCGA before using the more computationally demanding GMM.

Model selection for latent class models

Based on the number of classes identified in previous research, I applied 3 groups in a LCGA model initially and added more groups (up to a maximum of 6) in a step-wise manner. I compared models with linear growth (intercept and slope) and non-linear growth (intercept, linear, quadratic and cubic parameters) in turn. All LCGA estimations were run with 500 random starting value sets to avoid local solutions, a problem described by Jung and Wickrama as occurring:
“where during curve estimation a largest value (maximum) or smallest value (minimum) that a function takes is identified for only a given area on that curve, but that is not necessarily the largest or smallest value for the entire curve (i.e., the global minimum or maximum)”.

I fitted GMM models in a similar step-wise fashion, but used the estimated class-specific growth factor (e.g. intercept, slope and polynomial) means from the LCGA models (with the same number of classes and growth factors) as starting values. This is one of two strategies offered by Muthén and Muthén to try to bypass the common problem of convergence in GMM and, thus, find global maximum solutions. The other strategy is to use the estimated class-specific growth coefficient means and standard deviations from a $k$ class GMM model to compute starting values for a $k+1$ class model (by, for example, adding half the standard deviation to the mean).

I used the Bayesian information criterion values (BIC; a lower score indicating better model fit) and the bootstrapped likelihood ratio test (BLRT; obtained using the TECH14 option in Mplus) to guide model selection. In simulation studies, the BIC and BLRT have been shown to be the two best performing indices/tests to determine the number of classes in mixture modelling. The null hypothesis of the BLRT is that the model with $k-1$ classes fits the data as well as the model with $k$ classes, thus a low $p$-value points towards the $k$ class solution being a better fit of the model. I also required the final model to have latent classes that included at least 1% of the sample and an entropy value $>0.8$. A lower entropy value implies poorer separation of classes which, in this case, indicates that there is uncertainty around membership of trajectory groups.

To choose the final model, I compared the results of the final LCGA and GMM fitted models with the same number of classes and shape using the log likelihood values, with a higher value indicating a better fit. For the final selected model, I produced a plot showing the probability of symptoms at each time point for each trajectory group based on posterior distributions of asthma/wheeze prevalence. An example of Mplus code used to derive asthma/wheeze trajectories groups is included in Appendix 7.2.

7.3.3. Risk factors by asthma/wheeze trajectory group (objective 7B)

Firstly, I described cohort characteristics and missing data for risk factors using descriptive statistics (independent of the latent growth curve modelling described above). The association between variables and the probability of missingness in at least one study variable was
explored using multivariable logistic regression. There was a relatively high level of missing data across the cohort (10.3% of children had a missing value for at least one risk factor) and, prominently, I found that missing data was strongly associated with a particular birth hospital. Therefore, I imputed missing data before carrying out further analyses in this study.

**Multiple imputation**

Multiple imputation by chained equations (MICE) was used to impute missing values in risk factors for the multivariable logistic regression models. MICE generates imputed values using imputation models for each variable with missing values specified as a function of all other variables involved in the model of interest for the outcome and any other additional variable that may influence why values are missing. This approach allows the handling of different variable types (continuous, binary, unordered categorical and ordered categorical) that may be affected by missingness.\(^{452}\) It operates under the assumption that data are missing at random, conditional on the variables included in the imputation models. In other words, the probability that a value is missing depends on the observed variables that are being conditioned on in the imputation.\(^{453}\)

To meet this assumption I conditioned on all covariates used in the multivariable logistic regression models (introduced below), as well as: birth at the identified hospital with high levels of missingness (yes or no), NS-SEC, preterm-related complication (yes or no) and postnatal intensive care stay (yes or no). A binary indicator of any asthma/wheeze symptoms between age 2 and 9 was also included. Partially observed covariates were imputed using linear regression (for birthweight, gestational age, maternal age and number of older siblings), logistic regression (for severe maternal asthma and smoking status) and multivariable logistic regression (for number of older siblings and delivery method) models. Preterm birth, maternal age groups, siblings groups and SGA were defined using the relevant variables after imputation. Stata’s *mice* command was used to implement MICE in Stata 15.\(^{314}\)

**Characteristics of children by asthma/wheeze trajectory group**

Firstly, the distribution of risk factors, including missing data, by each child’s most likely asthma/wheeze trajectory group were described in percentages. I then used multinomial logistic regression on the imputed datasets to estimate the probability of belonging to a particular asthma/wheeze trajectory group, relative to the never or infrequent group, based on these risk factors. These analyses are all univariable (i.e. unadjusted for other risk factors). This is because rather than a prediction model, I wanted to describe the most likely composition of the identified trajectory groups. Further, these variables are so closely
correlated that any understanding of possible causal relationships would require a different focus in my analyses and a new investigation of the plausible pathways (including a new DAG).

I report results of the multinomial logistic regression models as estimated relative risk ratios (RRRs), which are the ratios of the relative probability of being in a trajectory group over the probability of being in the never or infrequent asthma/wheeze group. To account for error introduced through potential misclassification of children into trajectory groups via the latent class models, I employed what is known as the refined three-step procedure. This refined three-step procedure is an improvement to the standard three-step procedure often used with latent class modelling because it uses the probability of each individual being a member of all classes rather than just their most likely class; it has been shown to lead to less biased results than simpler three-step techniques. Univariable multivariate logistic regression analyses were replicated on all imputed datasets and the final estimates derived using Rubin's rules (implemented in Mplus). An example of Mplus code used to implement the multivariate logistic regression is included in Appendix 7.2.

7.3.4. Mediation analysis (objective 7C)

Counterfactual disparity measures

Drawing from my work in chapter 6, I will use the CDM to capture the proportion of socioeconomic disparity in asthma/wheeze that would remain if severe (i.e. hospitalisation for) bronchiolitis were intervened upon in infancy. For this analysis, I created a binary outcome indicating chronic asthma from the trajectory groups derived through LCGA. Cohort members with early-persistent and intermediate-onset asthma/wheeze as their most likely trajectory group in LCGA analyses were defined as having chronic asthma, and early-transient and no or infrequent class members were defined as not having chronic asthma. I initially tried to run this analysis using a categorical outcome (i.e. the four trajectory groups), but encountered problems of stability of the estimates due to the small numbers in some subgroups. I focussed on chronic asthma (as opposed to any asthma/wheeze) to delineate between symptoms that may be indicative of early wheeze rather than asthma.

I reformulated objective 7C to reflect the estimate of mediation that can be derived using the CDM estimand and the new binary outcome. The new objective was “to establish the magnitude of socioeconomic disparity in the presence of chronic asthma trajectory groups that would remain if no child had an admission for bronchiolitis during infancy”. This was defined as: \[ \text{CDM}(m=0) = \mathbb{E}[Y(0)|A=a] - \mathbb{E}[Y(0)|A=a^*], \] where \( A = \text{SIMD group} \) (a if high/medium, \( a^* \) if low),
where $M$ is $\geq 1$ hospital admission for bronchiolitis in the first year of life (1 if yes, 0 if no) and $Y$ is chronic asthma trajectory group (1 is yes, 0 if no). This captures the contrast between the potential risks of chronic asthma under high/medium compared to low levels of socioeconomic deprivation, were no infants to have a hospital admission for bronchiolitis in infancy.

As explained in Chapter 6, the assumptions usually required to identify and estimate causal effects from observational data are no interference, consistency, positivity and conditional exchangeability. In the context of both types of CDMs, the assumption of no interference would mean that one child’s potential outcome (of asthma/wheeze) was dependent on another child’s level of SEP, or another child’s hospitalization.\textsuperscript{380,387} Given that the causes of asthma are complex and manifold, but mostly dependent on one’s own levels of exposure, it is unlikely that interference is violated in this study. Consistency, another condition that is usually invoked to identify causal effects, requires the study exposure to be well-defined. However, as outlined in Section 6.2.1, this assumption is not required to be upheld when estimating the CDM. Through this approach we accept that the association between exposure and outcome carry “relevant disparity-related (but not necessarily causal) information”\textsuperscript{370}

With regards to the positivity assumption, this would require that within each stratum defined by the mediator-outcome confounders there is a non-zero probability that a child is hospitalised for bronchiolitis. Based on the large number of children in this study and small number of strata (particularly with a binary outcome) it is justified to assume that the positivity assumption is upheld.\textsuperscript{380} Through my study DAG for the relationship between bronchiolitis admission and asthma/wheeze group (see Figure 7.2), I have outlined the many confounders required for the conditional exchangeability assumption (i.e. exchangeable conditional on a set of covariates $C$) to be met. Some of these covariates are unavailable in the study dataset, and the implications of this on the conditional exchangeability assumption will be considered in the discussion.

**Inverse probability weighting of marginal structural models**

I used IPW of MSMs to estimate $\text{CDM}(m=0)$ for children in high and medium SIMD groups, compared to the low (deprivation) SIMD group. The steps to this approach, outlined in Box 6.2, began by generating weights from the exposure model using multinomial logistic regression and from the mediator model using logistic regression. I then fitted a weighted regression model of the outcome against the exposure, the mediator and the interaction between the exposure and mediator. The estimated coefficient for the mediator in this model represents the estimated $\text{CDM}(m=0)$. Area of residence, year and maternal country of birth were included in the propensity score models that generated the weights as confounders of
the exposure-outcome relationship. Area of residence, birth season birth year, chronic condition, delivery method, gestational age, maternal age, severe maternal asthma, maternal country of birth, maternal smoking, number of older siblings, SGA and child sex were included as confounders of the mediator-outcome relationship. Birthweight, which was available in the dataset and identified as a potential confounder, was not included due to its high correlation with gestational age and SGA.

The estimated $\text{CDM}(m=0)$ is displayed as the risk difference between high/medium and low SIMD groups in the risk of chronic asthma in childhood per 1,000 children if no child had hospital admission for bronchiolitis during infancy. I also quantified the absolute risk and risk difference of chronic asthma by SIMD group and proportion of this inequity explained by bronchiolitis admissions. As discussed in Chapter 6, SEP is not suitable for causal investigations as such, so I quantified the extent of the exposure-outcome association by estimating the marginal risk of chronic asthma by SIMD group using a logistic regression model of the outcome against the exposure using IPW. The weights used for this are found by logistic regression modelling of the exposure that includes the exposure-outcome confounders, year of birth, maternal country of birth and area of residence. As in Chapter 6, the proportion of inequity explained by bronchiolitis admissions was estimated as: $(\text{risk difference}-\text{CDM}(m=0))/\text{risk difference}\times 100$.

**Bootstrap inference method**

To calculate confidence intervals for the above estimates of $\text{CDM}(m=0)$, TCE and proportion of disparity explained, whilst addressing the presence of missing data in my dataset, I used single stochastic imputation using chained equations with 10 burn-in iterations as outlined by Micali et al.$^{457}$ Imputation was directly followed by the mediation analysis and both processes were repeated on 1000 bootstrap samples to estimate standard errors. I carried out the imputation under the assumption that missingness was at random, conditional on the variables included in the imputation models. To meet this assumption I conditioned on all covariates from the substantive model, alongside birth at the identified hospital with high levels of missingness (yes or no), NS-SEC, preterm-related complication (yes or no) and postnatal intensive care stay (yes or no). Partially observed covariates were imputed using linear regression (for birthweight, gestational age, maternal age and number of older siblings), logistic regression (for severe maternal asthma and smoking status) and multivariable logistic regression (for number of older siblings and delivery method) models. Following each imputation, preterm birth, maternal age groups, siblings groups and SGA were defined using the relevant variables. The Stata code for this analyses is available in Appendix 7.3.
7.4. Results

7.4.1. Missing data

As displayed in the flow chart (Figure 7.3), 2.7% of the original cohort of births between 1 January 2007 and 31 June 2008 were dropped from the dataset based on the predefined exclusion criteria. Of the remaining 83,853 children in the cohort, 10.3% (8,632) children had missing data for at least one variable. Missingness was driven by infants without linked maternal records (n=4,353, 5.2% of the cohort) and missing data on smoking status during pregnancy, which was missing for an additional 3,807 of the infants who were linked to mother’s records (9.7% missing in total cohort).

As shown in Table 7.3, missingness in any variable was strongly associated with area: 45.9% of children with missing data had a Glasgow postcode area. Further investigation revealed that one hospital in Glasgow in particular contained a high proportion of the missing information on maternal smoking status and I, therefore, included this hospital as a binary predictor when performing imputations. In multivariable logistic regression, infants born at this hospital had a 9-fold greater odds of being in the missing data group compared to the non-missing data group (aOR 8.99, 95% CI 8.33 to 9.70). Other notable predictors of missingness were: earlier birth year, spring birth, postal areas not in northern Scotland, mother born outside the UK and intensive care stay after birth. Based on the proportion of missingness in the dataset, 11 imputation datasets were generated through MICE.

---

**Figure 7.3.** Flow diagram showing derivation of final study cohort
Table 7.3. Characteristics of final study cohort (pre-imputation), number of children with missing data and estimated adjusted odds ratios (aORs) for missingness

<table>
<thead>
<tr>
<th></th>
<th>Whole cohort</th>
<th>%</th>
<th>Missing data on ≥1 risk factor</th>
<th>N</th>
<th>%</th>
<th>aOR** (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Total</strong></td>
<td>83,853</td>
<td>100.0</td>
<td>8,632</td>
<td>100.0</td>
<td>0.92</td>
<td>(0.80–1.07)</td>
</tr>
<tr>
<td><strong>SIMD</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Low</td>
<td>21,991</td>
<td>26.2</td>
<td>2,115</td>
<td>24.5</td>
<td>Reference</td>
<td></td>
</tr>
<tr>
<td>Medium</td>
<td>31,738</td>
<td>37.8</td>
<td>2,974</td>
<td>34.5</td>
<td>1.03</td>
<td>(0.97–1.10)</td>
</tr>
<tr>
<td>High</td>
<td>30,124</td>
<td>35.9</td>
<td>3,543</td>
<td>41.0</td>
<td>0.97</td>
<td>(0.90–1.04)</td>
</tr>
<tr>
<td><strong>Any asthma/wheeze</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>No</td>
<td>76,034</td>
<td>90.7</td>
<td>7,819</td>
<td>90.6</td>
<td>Reference</td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td>7,819</td>
<td>9.3</td>
<td>813</td>
<td>9.4</td>
<td>0.99</td>
<td>(0.92–1.08)</td>
</tr>
<tr>
<td><strong>Bronchiolitis</strong></td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>No</td>
<td>81,178</td>
<td>96.8</td>
<td>8,404</td>
<td>97.4</td>
<td>Reference</td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td>2,675</td>
<td>3.2</td>
<td>228</td>
<td>2.6</td>
<td>0.92</td>
<td>(0.80–1.07)</td>
</tr>
<tr>
<td><strong>Birth season</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Spring</td>
<td>27,967</td>
<td>33.4</td>
<td>3,045</td>
<td>35.3</td>
<td>Reference</td>
<td></td>
</tr>
<tr>
<td>Summer</td>
<td>19,180</td>
<td>22.9</td>
<td>1,787</td>
<td>20.7</td>
<td>0.80</td>
<td>(0.75–0.86)</td>
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<tr>
<td>Autumn</td>
<td>14,083</td>
<td>16.8</td>
<td>1,479</td>
<td>17.1</td>
<td>0.90</td>
<td>(0.83–0.96)</td>
</tr>
<tr>
<td>Winter</td>
<td>22,623</td>
<td>27.0</td>
<td>2,321</td>
<td>26.9</td>
<td>0.91</td>
<td>(0.85–0.96)</td>
</tr>
<tr>
<td><strong>Sex</strong></td>
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<td></td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>Female</td>
<td>40,972</td>
<td>48.9</td>
<td>4,212</td>
<td>48.8</td>
<td>Reference</td>
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<tr>
<td>Male</td>
<td>42,881</td>
<td>51.1</td>
<td>4,420</td>
<td>51.2</td>
<td>0.99</td>
<td>(0.95–1.04)</td>
</tr>
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<td><strong>Chronic</strong></td>
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<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>No</td>
<td>80,498</td>
<td>96.0</td>
<td>8,236</td>
<td>95.4</td>
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<td></td>
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<tr>
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<td>3,355</td>
<td>4.0</td>
<td>396</td>
<td>4.6</td>
<td>1.26</td>
<td>(1.13–1.41)</td>
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<td><strong>Area</strong></td>
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<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>Northern</td>
<td>10,903</td>
<td>13.0</td>
<td>566</td>
<td>6.6</td>
<td>Reference</td>
<td></td>
</tr>
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<td>Central</td>
<td>23,785</td>
<td>28.4</td>
<td>1,865</td>
<td>21.6</td>
<td>1.52</td>
<td>(1.37–1.67)</td>
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<td>Glasgow</td>
<td>19,516</td>
<td>23.3</td>
<td>3,966</td>
<td>45.9</td>
<td>1.42</td>
<td>(1.28–1.58)</td>
</tr>
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<td>Edinburgh</td>
<td>14,254</td>
<td>17.0</td>
<td>1,144</td>
<td>13.3</td>
<td>1.55</td>
<td>(1.39–1.72)</td>
</tr>
<tr>
<td>Southern</td>
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<td>18.4</td>
<td>1,091</td>
<td>12.6</td>
<td>1.22</td>
<td>(1.10–1.36)</td>
</tr>
<tr>
<td><strong>Maternal county of birth</strong></td>
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<td></td>
<td></td>
<td></td>
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</tr>
<tr>
<td>UK</td>
<td>74,792</td>
<td>89.2</td>
<td>7,579</td>
<td>87.8</td>
<td>0.88</td>
<td>(0.82–0.95)</td>
</tr>
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<td>Non-UK</td>
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<td>10.8</td>
<td>1,053</td>
<td>12.2</td>
<td>Reference</td>
<td></td>
</tr>
<tr>
<td><strong>Maternal age</strong></td>
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<td></td>
<td></td>
<td></td>
<td></td>
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</tr>
<tr>
<td>&lt;20</td>
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<td>7.2</td>
<td>560</td>
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<td>20-29</td>
<td>36,099</td>
<td>43.1</td>
<td>1,970</td>
<td>22.8</td>
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<td>35-39</td>
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<td>39.4</td>
<td>1,523</td>
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<td>&gt;40</td>
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<td>4,353</td>
<td>50.4</td>
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<td></td>
</tr>
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<td>2.4</td>
<td>102</td>
<td>1.2</td>
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<td>5.2</td>
<td>4,353</td>
<td>50.4</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Maternal smoking</strong></td>
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<td></td>
<td></td>
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<td></td>
</tr>
<tr>
<td>No</td>
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<td>70.5</td>
<td>371</td>
<td>4.3</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td>16,600</td>
<td>19.8</td>
<td>101</td>
<td>1.2</td>
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<tr>
<td>Missing</td>
<td>8,160</td>
<td>9.7</td>
<td>8,160</td>
<td>94.5</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Siblings</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>0</td>
<td>37,158</td>
<td>44.3</td>
<td>2,869</td>
<td>33.2</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1</td>
<td>26,701</td>
<td>31.8</td>
<td>863</td>
<td>10.0</td>
<td></td>
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</tr>
<tr>
<td>2+</td>
<td>15,337</td>
<td>18.3</td>
<td>243</td>
<td>2.8</td>
<td></td>
<td></td>
</tr>
<tr>
<td>missing</td>
<td>4,657</td>
<td>5.6</td>
<td>4,657</td>
<td>54.0</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Delivery method</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Vaginal</td>
<td>59,403</td>
<td>70.8</td>
<td>2,951</td>
<td>34.2</td>
<td></td>
<td></td>
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<tr>
<td>Elective caesarean</td>
<td>8,202</td>
<td>9.7</td>
<td>425</td>
<td>4.9</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Emergency caesarean</td>
<td>11,893</td>
<td>14.2</td>
<td>901</td>
<td>10.4</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Missing</td>
<td>4,355</td>
<td>5.2</td>
<td>4,355</td>
<td>50.5</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Premature birth</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>No</td>
<td>74,302</td>
<td>88.6</td>
<td>3,820</td>
<td>44.3</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td>5,140</td>
<td>6.1</td>
<td>401</td>
<td>4.6</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Missing</td>
<td>4,411</td>
<td>5.3</td>
<td>4,411</td>
<td>51.1</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>SGA</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>No</td>
<td>71,596</td>
<td>85.4</td>
<td>3,330</td>
<td>38.6</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td>7,442</td>
<td>8.9</td>
<td>487</td>
<td>5.6</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Missing</td>
<td>4,815</td>
<td>5.7</td>
<td>4,815</td>
<td>55.8</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
*Children meeting any asthma/wheeze definition between 2 and 9 years; **(aOR = adjusted odds ratio) multivariable logistic regression comparing the likelihood of being in the missing data group compared to the non-missing data group. Also included in the missingness model and, for simplification, not shown in table: birth at hospital with high levels of missing data, aOR 8.99 (95% CI 8.33 to 9.70); later birth year, aOR 0.87 (95% CI 0.82 to 0.92); intensive care stay, aOR 1.18 (95% CI 1.08 to 1.29); prematurity-related complication, aOR 1.01 (95% CI 0.87–1.18); and NS-SEC analytic classes.

7.4.2. Cohort characteristics

The final study cohort comprised 83,853 children of which 48.9% were female and 40.3% were registered with a Glasgow or Edinburgh postal area at birth (table 7.3). At least 6.1% infants were born before 37 weeks’ gestation and 8.9% were classified as SGA. Just under a quarter of births with a delivery method recorded were born via a caesarean section and 16,600 mothers (19.8%) reported smoking traditional cigarettes during their pregnancy (among those for whom delivery method and smoking data were available). Of the cohort, 0.3% (2,675) were admitted to hospital at least once with a diagnosis of bronchiolitis in the first year of life and 9.3% (7,819) met the study definition for asthma/wheeze at one time point or more between the ages 2 and 9. Information from the mother’s delivery record was available (i.e. linked) for 79,500 (94.8%) of children and the average follow-up time per child was 9.87 years. Of children in the cohort at age 2 years, 2,325 (2.8%) either emigrated (according to their CHI registry) or died by their 9th birthday (see Table 7.4).

Table 7.4 Observed completeness and frequency of asthma/wheeze symptoms, by age of child

<table>
<thead>
<tr>
<th>Age</th>
<th>Completeness of asthma/wheeze indicator</th>
<th>Asthma/wheeze symptoms N</th>
<th>% of cohort (N = 83,853)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Complete child record N</td>
<td>Child died or emigrated N</td>
<td>% of cohort (N = 83,853)</td>
</tr>
<tr>
<td>2</td>
<td>83,853</td>
<td>0</td>
<td>0.0%</td>
</tr>
<tr>
<td>3</td>
<td>83,298</td>
<td>555</td>
<td>0.7%</td>
</tr>
<tr>
<td>4</td>
<td>82,783</td>
<td>1,070</td>
<td>1.3%</td>
</tr>
<tr>
<td>5</td>
<td>82,431</td>
<td>1,422</td>
<td>1.7%</td>
</tr>
<tr>
<td>6</td>
<td>82,182</td>
<td>1,671</td>
<td>2.0%</td>
</tr>
<tr>
<td>7</td>
<td>81,959</td>
<td>1,894</td>
<td>2.3%</td>
</tr>
<tr>
<td>8</td>
<td>81,735</td>
<td>2,118</td>
<td>2.5%</td>
</tr>
<tr>
<td>9</td>
<td>81,528</td>
<td>2,325</td>
<td>2.8%</td>
</tr>
</tbody>
</table>
7.4.3. Growth modelling

Table 7.5 presents model fit indices for the LCGA and GMM growth modelling. Only results for two versions of GMM models are presented; the best fitting one is a 3-class linear model, which had a high entropy value (0.98) and a lower BIC value compared to the equivalent 3-class linear LCGA solution. However, a global maximum solution could not be obtained for this 3-class GMM model and one of its classes contained only 0.5% of the sample. GMM models with non-linear growth failed to converge, even after trying both strategies suggested by Muthén and Muthén (using estimated class-specific growth factors means/coefficients from LCGA/smaller GMM models) to determine starting values.

The LCGA model that identified 4 groups with cubic growth was selected as the most suitable model overall. This decision was guided by previous research, a low BIC value relative to other solutions, as well as groups each with >1% of the cohort (see Table 7.5). Figure 7.4 displays average latent class probabilities (classes defined by colour, see key), grouped by most likely class membership. The mean membership probabilities of classes in this 4-class cubic LCGA solution were between 0.89 and 0.99, and the overall entropy value was 0.97 indicating a well-classified model. The posterior distributions of asthma/wheeze prevalence at each time point by class are displayed in Figure 7.5.

The 4 trajectory groups are not necessarily true phenotypic groups, but represent descriptions of the variation in individual trajectories, where the prevalence of each group is based on estimated posterior probabilities. The groups can be summarised as:

1. Never or infrequent asthma/wheeze: 93.3% of the cohort had little to no asthma/wheeze symptoms at any time point.
2. Early-transient asthma/wheeze: 2.6% of the cohort had asthma/wheeze symptoms that had begun by age 2, had a peak prevalence at age 3 and dissipated by age 7.
3. Intermediate-onset asthma/wheeze: 2.2% of the cohort had asthma/wheeze symptoms that had begun by age 4 and had a peak prevalence at age 8, a year before follow-up ended.
4. Early-persistent asthma/wheeze: 1.8% of the cohort had asthma/wheeze symptoms that had begun by age 2, had a peak prevalence at age 6 and were ongoing at the end of follow-up.
Table 7.5. Model fit indices for LCGA and GMM on asthma/wheeze symptoms (selected model highlighted)

<table>
<thead>
<tr>
<th></th>
<th>LL</th>
<th>BIC</th>
<th>Entropy</th>
<th>VLMR (p)</th>
<th>BRLT (p)</th>
<th>Sample size per class (based on estimated posterior probabilities)</th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
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</thead>
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L=log likelihood, BIC=Bayesian Information Criteria, VLMR=Vuong-Lo-Mendell-Rubin likelihood ratio test, BLRT= bootstrap likelihood ratio test; *model did not reach global solution
Chapter 7 - Trajectories of asthma/wheeze

Figure 7.5. Average trajectory group probabilities*

*Classes are defined by colour (see key) and grouped by most likely class membership: results derived from the 4-class LCGA model with cubic growth.

Figure 7.4. Probability and 95% CI of symptoms at each time point for latent trajectory groups*

*Based on posterior distributions of asthma/wheeze prevalence, derived from the 4-class LCGA model with cubic growth. Never or infrequent asthma/wheeze group has a consistent near 0 probability of reporting asthma/wheeze symptoms.
7.4.1. Association between asthma/wheeze trajectories and risk factors

Table 7.6 and Figure 7.6 show the results of multinomial logistic regression models for class membership by asthma/wheeze risk factors. The results presented in this section are based on these results, which take into account error introduced through potential misclassification of children into trajectory groups. I have also included raw risk factor distribution by asthma/wheeze trajectories, including missing data, in Table A7.2 (Appendix 7.4). The results in this table are based on most likely class membership (i.e. not accounting for error) in the pre-imputation dataset. A notable take away from Table A7.2 is the difference in raw numbers and percentage of children with ≥1 hospital admission for bronchiolitis by trajectory group: 2.8% \((n = 2,206)\) in the never or infrequent group, 10.8% \((n=200)\) in the early-transient group, 9.0% \((n=139)\) in the early-persistent group and 5.5% \((n=90)\) in the intermediate-onset group.

As displayed in Figure 7.6, a graded effect of SIMD was observed across all asthma/wheeze groups relative to the never or infrequent group, with the most deprived SIMD group having the highest risk of asthma/wheeze. The most pronounced association was in the early-persistent group: children born into the most deprived SIMD group had a 65% greater relative risk \(\text{RRR 1.65, 95% CI 1.45 to 1.87}\) and children in the medium SIMD group had a 32% greater relative risk \(\text{RRR 1.32, 95% CI 1.15 to 1.50}\) of membership of this latent asthma/wheeze group than the never or infrequent group, compared to those born into the least deprived group. As illustrated in Figure 7.6, hospital admission for bronchiolitis and severe maternal asthma have the strongest unadjusted association with asthma/wheeze group membership. One or more hospital admission with a diagnosis of bronchiolitis in the first year of life increased the relative risk of early-transient asthma/wheeze group membership by 4.44 \((95\% \text{ CI 3.86 to 5.10})\), early-persistent asthma/wheeze group membership by 3.47 \((95\% \text{ CI 2.94 to 4.09})\) and intermediate-onset group membership by 1.83 \((95\% \text{ CI 1.47 to 2.28})\), compared to the never or infrequent group. Severe maternal asthma was most strongly associated with early-persistent asthma/wheeze \(\text{IRR 3.33, 95% CI 2.77 to 4.00}\), followed by the intermediate-onset group \(\text{RRR 2.71, 95% CI 2.22 to 3.31}\), then the early-transient group \(\text{RRR 1.96, 95% CI 1.56 to 2.45}\).

Other risk factors shown to have positive associations with group membership across all 3 asthma/wheeze groups, compared to no or infrequent asthma/wheeze, include: chronic condition \((46\%-95\% \text{ greater relative risk})\), male sex \((46\%-66\% \text{ greater relative risk})\); UK maternal country of birth \((22\%-30\% \text{ greater relative risk})\); maternal smoking during pregnancy \((23\%-37\% \text{ greater relative risk})\); emergency caesarean delivery method \((9\%-30\% \text{ greater relative risk compared to vaginal delivery})\); and SGA \((11\%-33\% \text{ greater relative risk})\). Notably, the strongest association for all these risk factors was with the early-persistent group. This was again the
case for preterm birth, which increased the relative risk of early-persistent group membership by 2.08 (95% CI 1.80 to 2.40), early-transient group membership by 1.90 (1.65 to 2.19) and intermediate-onset by 1.23 (95% CI 1.03 to 1.48). There was an S-shaped pattern to maternal age across asthma/wheeze groups, with slightly stronger associations in the youngest and oldest age groups. The presence of older siblings had a negatively graded association with the risk of membership of the early-transient and intermediate-onset groups and no clear association with early-persistent asthma/wheeze. A Glasgow or southern Scottish compared to an Edinburgh postal area at birth was associated with membership of the early-transient and early-persistent group.

Table 7.6. Univariable RRRs (95% CI) of belonging to asthma/wheeze groups compared to the never or infrequent asthma/wheeze group, by risk factor

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<th>Risk factor</th>
<th>Early-Transient</th>
<th>Early-Persistent</th>
<th>Intermediate-onset</th>
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<td>1.32 (1.15–1.50)</td>
<td>1.24 (1.10–1.41)</td>
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<td>1.65 (1.45–1.87)</td>
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<td>1.83 (1.47–2.28)</td>
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<td>Autumn</td>
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<td>Reference</td>
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<tr>
<td>Winter</td>
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<td>1.10 (0.94–1.27)</td>
<td>1.03 (0.89–1.19)</td>
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<td>1.13 (0.98–1.31)</td>
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<td>1.46 (1.33–1.60)</td>
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</tr>
<tr>
<td><strong>Caesarean delivery</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Elective vs. vaginal</td>
<td>0.98 (0.85–1.14)</td>
<td>1.10 (0.93–1.29)</td>
<td>1.06 (0.91–1.24)</td>
</tr>
<tr>
<td>Emergency vs. vaginal</td>
<td>1.18 (1.04–1.33)</td>
<td>1.30 (1.15–1.48)</td>
<td>1.09 (0.95–1.24)</td>
</tr>
<tr>
<td><strong>Preterm birth</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>yes vs. no</td>
<td>1.90 (1.65–2.19)</td>
<td>2.08 (1.80–2.40)</td>
<td>1.23 (1.03–1.48)</td>
</tr>
<tr>
<td><strong>SGA</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>yes vs. no</td>
<td>1.11 (0.96–1.29)</td>
<td>1.33 (1.15–1.54)</td>
<td>1.13 (0.97–1.31)</td>
</tr>
<tr>
<td><strong>Year</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2008 vs. 2007</td>
<td>0.91 (0.83–1.00)</td>
<td>0.99 (0.90–1.10)</td>
<td>0.96 (0.88–1.06)</td>
</tr>
</tbody>
</table>
Figure 7.6. Forest plots showing univariable RRR estimates with 95% CI for membership of asthma/wheeze groups compared to the never or infrequent asthma/wheeze group, by risk factor.
7.4.2. Mediation analysis

As shown in Table 7.7, after accounting for exposure-outcome confounders, the potential risk of chronic asthma by age 10 would be 30.8 per 1000 children (95% CI 28.3 to 33.3) if all children were assigned to the low SIMD group at birth. This compares to 37.7 per 1000 for the medium SIMD group (95% CI 36.2 to 39.3) and 43.7 per 1000 for the high SIMD group (95% CI 41.3 to 46.1). The $CDM(m=0)$, calculated using IPW of MSMs, suggests that if no children had a hospital admission for bronchiolitis in infancy, the counterfactual disparity between children in the high compared to low SIMD group would be reduced from 13.0 chronic asthma cases per 1000 children (95% CI 9.6 to 16.4) to 10.7 per 1000 (95% CI 7.3 to 14.0). This is an estimated disparity reduction of 17.9% (95% CI 10.4 to 25.4). A larger proportional disparity reduction was estimated between the medium and low SIMD groups (21.2%, 95% CI 4.9 to 37.5). This reflects a larger relative (but lower absolute) change in potential risk difference, from 7.0 chronic asthma cases per 1000 (95% 3.7 to 10.3) before to 5.5 per 1000 (95% CI 2.3 to 8.8) after counterfactual disparity in bronchiolitis admission is removed.

Table 7.7. Potential risk of chronic asthma* by age 9, estimated counterfactual disparity in risk and corresponding disparity reduction involving bronchiolitis admissions, by SIMD group*

<table>
<thead>
<tr>
<th>SIMD group</th>
<th>Marginal risk per 1000 children** (95% CI)</th>
<th>Marginal risk difference per 1000 children** (95% CI)</th>
<th>$CDM(m=0)$ per 1000 children (95% CI)†</th>
<th>% disparity reduction (95% CI)‡</th>
</tr>
</thead>
<tbody>
<tr>
<td>IPW MSMs</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Low</td>
<td>30.8 (38.3, 33.3)</td>
<td>Reference</td>
<td>Reference</td>
<td>Reference</td>
</tr>
<tr>
<td>Medium</td>
<td>37.7 (36.2, 39.3)</td>
<td>7.0 (3.7, 10.3)</td>
<td>5.5 (2.3, 8.8)</td>
<td>21.2 (4.9, 37.5)</td>
</tr>
<tr>
<td>High</td>
<td>43.7 (41.3, 46.1)</td>
<td>13.0 (9.6, 16.4)</td>
<td>10.7 (7.3, 14.0)</td>
<td>17.9 (10.4, 25.4)</td>
</tr>
</tbody>
</table>

*binary outcome created from most likely trajectory group membership (early-persistent and intermediate-onset indicating yes); ** Risk and absolute risk difference after accounting for differences in year of birth, maternal country of birth and area of residence between groups using IPW; †95% CIs calculated using the bootstrap with 1000 replications; ‡disparity reduction (%) was estimated as $= (\text{risk difference} - \text{CDM}(m=0)) / \text{risk difference} \times 100$

7.5. Discussion

Latent growth modelling applied to longitudinal observations of asthma and wheeze symptoms retrieved from administrative health records led to the identification of four different asthma/wheeze trajectories between ages 2 and 9 years. I labelled the groups as follows:
never or infrequent (93.3% of the cohort), early-transient (2.6%), early-persistent (2.2%), and intermediate-onset asthma/wheeze (1.8%) trajectory groups. There was a socioeconomic gradient to group membership across all asthma/wheeze groups compared to the never or infrequent group, with children in the most deprived group having between 42-65% increased risk of asthma/wheeze compared to children in the least deprived group. At least one admission to hospital with bronchiolitis was strongly associated with asthma/wheeze group membership, particularly the early-transient and early-persistent groups, which had a 4.4 times and 3.5 times greater risk respectively. I estimated that if bronchiolitis admissions in infancy were eliminated then up to 18% of the difference in the risk of chronic asthma between children from the most and least deprived 30% of SIMD would be removed.

7.5.1. Strengths and weaknesses

Using administrative data, this study was able to follow up more than 80,000 children until their tenth birthday to determine asthma/wheeze trajectories. I am aware of only one other study, based in Canada, that has modelled asthma trajectories on such a large population-based level. A range of clinical definitions were used to describe asthma/wheeze in this study with input from a clinician, in contrast to the commonly used self- or parental-reported measures used elsewhere. This approach, paired with a cautious methodology to defining symptoms by dispensed medication (≥4 prescriptions in a 12 month period), means there are likely to be less false positives captured in this study. This is mirrored by the high proportion of children classified as most likely belonging to the none or infrequent asthma/wheeze group: over 93% of the study population compared to 82-83% in studies using the Millennium Cohort Study.

On the other hand, some children (likely with milder symptoms) may have been misclassified as having no symptoms in this study. The addition of primary care records in future administrative data studies may improve identification of these cases. Using only indicators of asthma and wheeze presumes that only these latent variables underlie the heterogeneity in how these conditions evolve over time. Other studies have included measures of atopy in modelling of asthma trajectories; however, overlap between medication used to treat asthma and atopic conditions meant that the current dataset would not have allowed for this nuanced differentiation. In addition, although I used predetermined criteria to determine trajectories, these methods still have a degree of subjectivity. Other studies have invoked avoidance of classes <5% of the study population; however, this seemed unnecessarily restrictive in a sample where 5% of the sample is more than 4,000 children.

I had to take other several pragmatic steps in this study based on constraints of the datasets. I restricted the sample to children born between 2007 and mid-2008 to obtain a 10-year follow
up period that included PIS data (only available from 2009 onwards) from age 2 for all cohort members, which was necessary to model asthma outcomes. The lack of complete PIS data before 2009 meant that I could not use dispensed medication data to define maternal asthma prior to their child’s birth. I instead relied on diagnoses during hospital admissions, leading to an indicator that would have mostly captured more severe and poorly managed cases of asthma. Given that poorly controlled symptoms are more prevalent in the most deprived socioeconomic groups, this limited definition of maternal asthma likely conflates the association between SEP and maternal asthma. I dichotomised the outcome for the mediation analysis into a binary indicator of chronic asthma/wheeze. I considered examining the four classes, but encountered problems of stability of the estimates due to the small numbers in some subgroups. In the future, cohort studies from Scottish administrative data with births spanning several years will enable longer follow up periods and a greater sample size for more nuanced analyses.

LCGA, used to define trajectory groups in this study, assumes zero variance within classes and may not be the best statistical representation of the underlying population. Successive cubic LCGA models (with more classes) led to an increasingly improved fit according to BIC and BLRT values, and the 4-class solution was chosen because of small class sizes in the 5-class model. Berlin and colleagues write that, rather than signifying a better fitting model, the pattern towards increasing classes may actually be the result of a poorly-specified model in data that has substantial within-class variability.\textsuperscript{461} I also modelled trajectories in the data using GMM, which offers a more flexible approach to variability within classes. However, non-linear GMM models failed to converge—a known problem with these models.\textsuperscript{439,461} As my aim was to replicate classes found in previous studies within a Scottish context (and use these to look further at pathways to inequities in asthma), rather than produce new trajectories, the resemblance of classes in this study to those found in numerous other studies gives me confidence with the results presented.

I used a three-step approach to growth modelling, which meant that I firstly created the latent classes and then used these classes in further analyses. Asparouhov and Muthén write that one-step models, which combine latent class modelling with the secondary model, can be flawed as the additional variables may affect latent class formation and result in unintended changes to the latent class meaning.\textsuperscript{454} I included classification errors in analyses looking at the association between risk factors and trajectories to account for potential latent class misclassification introduced in latent class analysis.\textsuperscript{191} I was unable to do this in the mediation analysis, which has the potential to affect the precision of estimates, although may not be an issue in light of the well-defined classes in this study (see Figure 7.4).
There were several confounders identified in previous research that could not be included in this study, which may have led to an overestimation of the mediating effect of bronchiolitis. This includes breastfeeding, which has broadly shown to be a protective factor for bronchiolitis and asthma, and associated with lower levels of deprivation,\textsuperscript{194,196,197} and outdoor air pollution and damp/mould, which are positively associated with socioeconomic deprivation and respiratory ill-health.\textsuperscript{116,125,153,202} The confounding effect of some unmeasured confounders may be partially captured by other confounders included in the model. For example, passive smoke exposure after birth by maternal smoking, and ethnicity by maternal country of birth. The pathways with which the final three unmeasured confounders (gestational diabetes, pre-eclampsia and breech presentation birth) are thought to influence offspring respiratory are also included as confounders e.g. delivery method and preterm birth.\textsuperscript{182,462}

Future studies utilising administrative data could be enhanced with linkage to other datasets, for example the Department for Environments Food and Rural Affairs’ modelled background pollution data,\textsuperscript{463} which will enable more of these confounders to be accounted for and bias to be eliminated from calculations. Alternatively, purposefully designed cohort studies offer detailed information on many variables that are not typically available in administrative data sources. For example, the Millennium Cohort Study has been used to examine the complex causal pathways between socioeconomic circumstances and adolescent mental health.\textsuperscript{464}

7.5.2. Comparison to other studies

The classes identified in this study are similar to those found in multiple other studies, showing four trajectories with distinct timing in the onset and longevity of symptoms. As noted above, the size of the classes were smaller (proportionally) than those found in other studies; however, the relative size between asthma/wheeze trajectory groups is similar. I will now compare my results to those of other studies, drawing particularly from the 2020 systematic review and meta-analysis of childhood wheeze trajectory-specific risk factors by Owora and Zhang.\textsuperscript{197} These authors note that there are few risk factor associations that are trajectory specific across the literature (i.e. there are similar associations for all trajectory groups compared to the never or infrequent group), which is broadly the finding of my study.

\textit{Early-transient asthma/wheeze}

Asthma/wheeze symptoms that begin early, but cease after the first few years of life have been identified and defined as early transient wheeze across the literature.\textsuperscript{197} Membership of this group is strongly associated with viral respiratory infection. In my study, hospital admission for bronchiolitis in infancy was associated with a 4.4 times increased risk of early-transient
asthma/wheeze group membership. This trajectory group has been found to have a low risk of atopy and of future asthma, but to still exhibit diminished airway function in mid-childhood. Greater number of siblings was associated with a low relative risk of membership of this group in my study. In contrast, Owora and Zhang found a 47% greater odds of early-transient wheeze group membership among children with siblings in their meta-analysis (pooled OR 1.47, 95% CI 1.06 to 2.04). The exclusion of symptoms under 2 years in my study outcome, the ages where children are most sensitive to infections and thus the influence of virus transmission from older siblings, may contribute to these findings.

**Early persistent asthma/wheeze**

Prolonged asthma/wheeze symptoms that begin in early life are associated with the poorest prognosis in term of lung function out of the asthma groups identified in other research. In my study, membership of this group had the strongest association with almost all risk factors, including socioeconomic position, relative to other asthma/wheeze groups. This group appears to mirror non-atopic asthma groups found in other studies; however, as a measure of atopy was not captured in my study this cannot be confirmed. In a trajectory analysis of asthma and atopy symptoms using the Millennium Cohort Study, children who were in the poorest households at 9 months, had the highest likelihood of being in the group with high wheeze and no atopy at ages 3, 5 and 7. Conversely, the most advantaged group were most likely to be in the low wheeze but high atopy group. As in Owora and Zhang’s meta-analysis, family history was most predictive of early-persistent wheeze (maternal asthma was associated with more than 3-fold increased relative risk compared to no or infrequent asthma/wheeze in my study).

**Intermediate-onset asthma/wheeze**

Two asthma/wheeze trajectories that begin at different points in mid-childhood have been described in other studies, labelled intermediate-onset (beginning between 3 and 5 years) and late onset (beginning between 6 and 8 years) by Owora and Zhang. I identified only one of these groups my final 4-class model, perhaps owing to my relatively strict definition of asthma/wheeze and the lack of follow up beyond the age of ten. In future work with a longer follow up, differences in the number of identified classes by stringency of asthma/wheeze definition could be explored. In my analysis, the intermediate-onset group had broadly similar but weaker associations with the included risk factors compared to the early-transient and early-persistent groups (versus the never/infrequent group). This group appears to mirror an atopic asthma phenotype found in other studies, although this association cannot be ascertained from my study. Risk factors operating in early childhood (as opposed to the pre-
and perinatal risk factors included in my study, such as growing up in a non-farming household and a lack of pet ownership, may partially explain the development of later onset atopic childhood asthma/wheeze.\textsuperscript{430,466}

\textit{The contribution of bronchiolitis to chronic childhood asthma}

There has been a longstanding debate on the link between early RSV-related lower respiratory tract infections (including bronchiolitis) and the development of asthma, although recent research point towards a non-causal relationship.\textsuperscript{467} Two systematic reviews of the literature on the link between early RSV-related lower respiratory tract infections and subsequent pulmonary function were published in 2020.\textsuperscript{468,469} One of these concluded that the evidence was “not overwhelming with conflicting results between studies”.\textsuperscript{469} p.1567 The other review reached a firmer conclusion: “Our findings, limited to exposure and immunoprophylaxis studies, do not support basing policy decisions on an assumption that prevention of RSV-LRTI will reduce recurrent chronic wheezing illnesses.”\textsuperscript{468} p.795 Notably, both reviews identified a high risk of confounding bias in observational studies.\textsuperscript{468,469}

Novelly, I have used causal inference methods, with explicit discussions of the involved assumptions, to reach the estimates in the current observational study. This is the first time the pathways between socioeconomic position, bronchiolitis and asthma have been explored using these methods. In my study, I estimate that up to 18\% of the disparity between socioeconomic groups (at the extreme 30 percentiles of SIMD) may be attributable to hospital admission for bronchiolitis during infancy. Moreover, it shows that there is at least 82\% of the association unexplained by bronchiolitis and broadly highlights the need to further investigate the inequities in admissions using causal inference methods. Notably, as part of this work, I have recognised some key confounders that are missing from my analyses, including air pollution, a lack of breastfeeding and damp/mouldy housing conditions. Research including information on other identified confounders is needed to clarify these pathways further.

7.6. \textbf{Chapter summary}

In this chapter, I have used trajectory analysis methods to define groups of asthma/wheeze within a Scottish population. I identified four distinct trajectories of symptoms in early life, labelled no or infrequent, early-transient, early-persistent and intermediate-onset asthma/wheeze, which is similar to those found in other studies. I looked at the association of membership of these groups with several early life risk factors and found notable associations with bronchiolitis and severe maternal asthma. There was also a clear socioeconomic gradient
to group membership. Using counterfactual disparity measures, I quantified the role of bronchiolitis admissions in the relationship between socioeconomic position and chronic asthma. I estimated that up to 18% of the inequity between groups was explained by hospital admissions for bronchiolitis in infancy, although unmeasured confounding in my analyses limits the causal interpretation of this finding.
Chapter 8. Discussion

Recap and chapter overview

I began this thesis by presenting an overview of research in the area of socioeconomic inequity, bronchiolitis and asthma among paediatric populations in the UK. I outlined the administrative data sources that enabled me to conduct my research and discussed the emerging issue of data opt outs to research using these types of data sources. In total, I carried out four main studies to look at the association between socioeconomic deprivation and bronchiolitis and asthma, two common and highly burdensome respiratory conditions. This final chapter draws on information from all these chapters to highlight the overall findings, as well as the strengths and limitations of my thesis. I will highlight the unique contribution of my work, before outlining the main implications of the key findings on future policy, practice and research. Lastly, I will re-assess my study implications in the context of the SARS-CoV-2 pandemic, which was ongoing at the time of writing.

8.1. Critical assessment of my PhD

8.1.1. Restatement of thesis rationale and aim

Bronchiolitis and asthma are major causes of morbidity among children in the UK and are responsible for substantial burden on the national health care system.\textsuperscript{4,32} Despite this, there are currently no preventative or curative measures for the majority of children that develop bronchiolitis or asthma.\textsuperscript{15,47} There is a large body of evidence investigating environmental, social and biological risk factors for these conditions, but much of this work does not fully consider the ways in which various risk factors interact with each other (and potentially bias results). There is also a myriad of research looking specifically at the link between RSV bronchiolitis and subsequent asthma,\textsuperscript{53,56,264} but precise risk factors leading to both conditions remain elusive. What we do know is that a prevailing facet of both conditions is the inequity in prevalence across socioeconomic groups.\textsuperscript{4,29} Identifying the factors that may mediate or confound the pathway between socioeconomic deprivation, bronchiolitis and asthma will help to unravel this complex picture. This is a pertinent step to working out how and when to best intervene to prevent either or both of the conditions. By focussing on early life risk factors, the aim is to reduce the burden of bronchiolitis and asthma by closing the gap in the incidence/prevalence of these conditions between children born into the most and least deprived socioeconomic groups.
To guide the structure of my work, I drew from the WHO’s multifaceted conceptual framework developed for the Commission on Social Determinants of Health (CSDH). As displayed in Figure 8.1, this framework differentiates between the levels of factors that likely influence health outcomes (in this case, the incidence of bronchiolitis and prevalence of asthma). According to the CSDH, at the outmost level are the wider social, economic and political context of society, which, in turn, stratify the population into socioeconomic positions (SEP). I do not include this outer level in my diagram, choosing to instead focus my work on the factors that sit between SEP, bronchiolitis and asthma because these pathways arguably represent the more pragmatic and achievable approach to intervention. My results add to the substantial evidence base showing the early origins of socioeconomic inequities in disease, which highlights that a broader approach to tackling the unequal distribution of health outcomes is warranted.

I focused broadly on SEP as the structural determinant of health in my project rather than more specific strata, which can include income, education, occupation, ethnicity and gender. I bring in specific components of the physical and built environment that likely interact with SEP and then, in turn, influence the presence of intermediary factors to shape inequities in health outcomes. For bronchiolitis and asthma, intermediary factors operating in the pre-natal and early post-natal period include tobacco smoke and air pollution exposure, premature birth, chronic conditions and poor housing conditions (shown in Figure 8.1). My project focuses on specific components of this framework to better explain how inequities in bronchiolitis and asthma arise.
Specifically, the aim of this PhD was to describe associations and explore pathways between early life socioeconomic deprivation, infant bronchiolitis and asthma in mid-childhood.

Focussing on specific pathways involved careful consideration of covariates, statistical methods and related assumptions, which I clearly outlined in each study. Relatedly, the datasets for this work are national birth cohorts created from English and Scottish administrative health and vital event datasets that, although extremely beneficial to epidemiological analyses, bring their own set of complications. Underlying my project aim are four major objectives, and I will now outline the key findings for each in turn.

8.1.2. Key findings

**Thesis objective 1** To describe the association between seasonality, socioeconomic position and bronchiolitis admission rates among infants across England

*Local area-level (Chapter 4)*

- There was a 5.3-fold difference in rates of bronchiolitis admissions across Clinical Commissioning Groups (CCGs) in England, with about one quarter of the variation explained in rates by area-level IMD scores
- The peak epidemic week of hospital admissions for bronchiolitis varied by 3 weeks across CCGs and tended to be earlier among CCGs with higher population densities, such as those located around/within London and Manchester

*Individual level (Chapter 5)*

- The seasonality of admissions varied marginally by individual-level socioeconomic group; the predicted peak epidemic week was 49.4 among infants in the least deprived IMD quintile and 50.3 in the most deprived IMD quintile
- Socioeconomic disparities in admissions to hospital for bronchiolitis persisted after taking seasonality into account, with a 40% greater risk of admissions among infants in highest compared to lowest quintile of IMD at week 50 (peak admission week)

**Thesis objective 2** To investigate the mediating contribution of maternal smoking in the pathway between early life socioeconomic position and bronchiolitis admission rates in infancy (Chapter 6)
Chapter 8 - Discussion

- Among children born in Scotland between 2012 and 2016, I estimated that the potential risk of bronchiolitis hospital admission in infancy would have been 57.1 per 1000 live births if all children were in the most deprived 30% of SIMD at birth.
- This was 16.3 more children with admissions for bronchiolitis for every 1000 infants were all children in the least deprived 30% of SIMD at birth.
- Using two methods to calculate the counterfactual disparity measure, I estimated that the elimination of maternal smoking would reduce about one fifth of the disparity in the proportion of children who had at least one hospital admission for bronchiolitis in the first year of life.

Thesis objective 3 To describe typical trajectories of asthma/wheeze among children <10 years in Scotland and their association with early life risk factors, including SEP (Chapter 7)

- I identified four trajectories of asthma/wheeze symptoms from the Scottish dataset, which I labelled never or infrequent (93.3% of the cohort), early-transient (2.6%), early-persistent (2.2%), and intermediate-onset asthma/wheeze (1.8%).
- There was a socioeconomic gradient to group membership across all asthma/wheeze groups compared to the never/infrequent group, with children in the most deprived 30% of SIMD having between 42-65% increased risk of asthma/wheeze compared to children in the least deprived 30% of SIMD.
- At least once hospital admission for bronchiolitis in infancy was strongly associated with asthma/wheeze group membership in unadjusted analyses, particularly the early-transient and early-persistent groups, which had a 4.4 times and 3.5 times greater risk respectively.

Thesis objective 4 To investigate the mediating contribution of bronchiolitis in the pathway between early life SEP and mid-childhood asthma/wheeze trajectory groups (Chapter 7)

- Among infants born in Scotland between January 2007 and June 2008, I estimated that the prevalence of chronic asthma by age 9 would have been 4.4% if all children were in the most deprived 30% of SIMD at birth compared to 3.1% if all children were in the least deprived 30% of SIMD at birth.
- I estimated that the elimination of bronchiolitis requiring hospital admission could reduce up to 18% of the disparity in the proportion of children with chronic asthma.
- I identified air pollution, housing conditions and breastfeeding as key risk factors to be incorporated into future research utilising similar causal inference methods.
8.1.3. Strengths and limitations

I have discussed the strengths and limitations specific to each study within the relevant chapter. Here, I bring together some of the major themes running through this thesis. Namely, the use of administrative data sources, the measurement of my exposure (SEP) and outcomes (bronchiolitis and asthma), and the statistical methods employed to address my thesis objectives.

**Administrative data sources**

Using administrative data sources in my thesis meant that I was able to follow up almost the whole population of children born in England and Scotland over the specified years of study. These large datasets (approximately 620,000 births in England and 53,000 in Scotland per year) meant that I could explore complex pathways that included relatively rare events and subgroups of the population. There have been very few studies that have looked at the mechanisms of the seasonality of RSV bronchiolitis using epidemiological methods, and none in a UK setting. Using nationwide hospital admission data in England, I was able to do this for bronchiolitis (a common but distressing condition) in a vulnerable population (infants), whilst focussing on the pressing issue of socioeconomic inequities. I created a Scottish birth cohort with 9 years of follow-up, enabling the exploration of longitudinal patterns of asthma/wheeze symptoms identified using objective measures.

However, I also encountered several drawbacks to using these data sources in my research. This was particularly apparent for the Scottish dataset as I was involved in the process of obtaining, cleaning and using the data from the beginning to end. Obtaining access to this data was a complex and time intensive process (taking 13 months to receive the data following approval of the data request). As part of the data request, we had to specify the exact variables needed for analyses and the reasons why. This meant that I needed to know the desired variables for analysis at the beginning of the process (meaning some later identified clinical risk factors, for example, gestational diabetes and pre-eclampsia, were not available to me). There were issues with the completeness of data. As shown in Chapter 6, patient records from one hospital in Glasgow in particular had a 9-fold greater odds of missing data. I also found that the completion of ethnicity information, a factor that is integral to fully understanding disparities in health outcomes, to be particularly poor (although, improving over time). There are wider ethical issues to the use of these datasets (also, for the most part, beyond the influence of researchers) and I identified patient opt-outs as a particular problem in my work.

**Socioeconomic position**
In Chapter 1, Section 1.2.2, I write about the benefits and disadvantages of IMD (the measure of SEP in my research) at some length. Listed benefits include its national applicability, composite construction and broad use across other health inequity research (aiding comparability of results). However, I also describe the potential issue of mathematical coupling in health inequity research where IMD (composed of 7 domains, one of which is health based) is used as the exposure. I investigated this issue in Chapter 5 by replicating my analysis with the health deprivation and disability domain excluded from IMD and found only slight differences in my results, giving me confidence that this did not bias my inferences. The components within the health domain are all based on adult health (including premature mortality and hospital admissions in adults) and therefore may still pose greater issues when looking at health outcomes in non-paediatric populations.

Another problem identified in Chapter 1 with using IMD as an indicator of SEP is its lack of specificity in comparison to other measures. This issue was revisited in Chapter 6 in regards the consistency assumption for causal inference. Here, I described how the exposure definition must have enough precision that any variation in that exposure does not lead to a different outcome. There was another measure of SEP available in the Scottish dataset based on occupation, the UK National Statistics Socio-Economic Classification (NS-SEC), which may have been better suited to meet the assumption of consistency. However, occupational categorisations like NS-SEC have been criticised for not fully capturing contemporary occupational structures, particularly given its basis of male-orientated job classifications. Instead, I used a method of causal inference (the counterfactual disparity measures (CDMs)), which requires less stringent assumptions for estimation and have been careful in the interpretation of my findings.

Defining bronchiolitis and asthma

Both health outcomes studied in this thesis rely on subjective clinical judgement for diagnosis, so a certain amount of imprecision in their identification is likely. The use of clinical definitions are particularly advantageous in comparison to parent-reported cases, which may be influenced by recall biases. I identified bronchiolitis through ICD-10 diagnosis codes in hospital admission records, limited to admissions where the patient was less than 1 year of age (as 90% of admissions for this condition occur in infancy). I did not have information on other (non-hospital admission) health care contacts for bronchiolitis, which means the captured cases likely represent the most severe cases of infection. It also means that capturing bronchiolitis cases is dependent on admission thresholds, highlighted in Chapter 4 where London was shown to have far lower rates of admissions (likely stemming from higher threshold for admission from accident and emergency departments). Alongside
admission and death records, I used dispensed medication information to define asthma/wheeze (with clinical input). Asthma is an ambulatory care sensitive condition, meaning that it ought to be preventable by interventions in primary care. Reliance on hospital admission records for this outcome would have biased my research in favour of children from lower socioeconomic groups, who are far more likely to have emergency treatment for this condition.

8.2. Implications of my research findings

8.2.1. Continuing high and unequal burden of disease

As outlined in Section 1.8.3 (Chapter 1), respiratory diseases, particularly asthma, have been identified as an important focus of public health policy in England and Scotland. My findings add weight to the need to focus resources on preventing both bronchiolitis and asthma as particularly burdensome conditions in the paediatric population. I showed that between 2012 and 2016 almost four in every ten infant admissions to hospitals in England during December included a diagnosis of bronchiolitis. More than 9% of children born between 2007 and mid-2008 in Scotland had symptoms of asthma and/or wheeze at least once that were severe enough to meet my stringent definition.

Prominently, I identified a socioeconomic gradient to the incidence of bronchiolitis admissions, and in the prevalence of all three identified trajectories of asthma/wheeze. This is not a new finding, although the detailed identification inequities across three asthma trajectories in a Scottish context is. Policies in England and Scotland have clearly highlighted the role of socioeconomic inequities in the development of respiratory ill health. The NHS England Long Term Plan, published in 2019, specifically refers to the higher incidence of respiratory disease in “disadvantaged groups and areas of social deprivation”. However, my findings highlight the importance of retaining focus on the social determinants of health and applying interventions in a way that brings down this gradient (i.e. proportionate universalism). This is particularly pertinent in restating in the context of asthma, for which policy has traditionally focused on controlling severe symptoms and reducing hospital admissions, rather than the prevention of disease.

8.2.2. Clinical practice and policy

By focusing on particular aspects of the framework in Figure 8.1, my work begins to untangle some of the ways in which preventative measures and interventions might be best applied to alleviate the burden of bronchiolitis and asthma.
Prevention of respiratory syncytial virus (RSV) infections

The first two studies in my thesis looked at the interplay of socioeconomic position and wider ecological factors on rates of hospital admissions for bronchiolitis (the left hand side of the framework displayed in Figure 8.1). Patterns of admissions can be viewed as a proxy for the circulation of RSV, which has been shown to cause the vast majority of bronchiolitis cases that lead to hospitalisation.\(^6\) I found differences in the timing of the epidemic peak at the CCG-level. The peak tended to occur earlier in CCGs that have higher population densities and are within/near major cities, such as London and Manchester. This highlights the potential role of population mixing, and possibly international travel, in the early spread of RSV. These findings have implications on preparedness for bronchiolitis infections. An early warning system could be set up for hospitals in these cities to alert Public Health England and the NHS at the annual onset of admissions for bronchiolitis. These systems could trigger surge planning in paediatric departments in other hospitals, inform the distribution of prophylaxis or future vaccinations, and be used to time simple public health messages, such as warning families to avoid taking their young infants into public spaces or encourage frequent handwashing for older siblings.

I found no clear differences in the timing of admissions for bronchiolitis by SEP at the individual level. This suggests that differential timing of administering palivizumab prophylaxis—the one currently available preventative measure\(^3\)—may not be the best course of action to tackle inequities in the incidence of this condition. It also suggests that information campaigns to reduce the spread of RSV, such as handwashing, are also unlikely to reduce inequities (as would perhaps be the case had I found that RSV spreads quicker among more socioeconomically deprived families). Instead, I found continued higher levels of admissions for bronchiolitis among infants born into the most socioeconomically deprived households, even after accounting for seasonal patterns. This presents evidence that to reduce inequities in admissions for bronchiolitis, the focus should be on tackling the chronic exposures that make children in more deprived groups susceptible to severe illness when infected with RSV (see below).

It is likely that a vaccine for RSV for pregnant women, which aims to protect infants during the first few months of life, will become available within the next 5-10 years.\(^42\) The evidence of unequal uptake of vaccines for children across socioeconomic groups,\(^247\) paired with a greater reluctance for vaccination among pregnant women,\(^244\) means that a vaccine for RSV in pregnancy has the potential to widen inequities in rates of bronchiolitis further. Embedding maternal vaccination within routine antenatal care (as opposed to a separate visit to a primary care centre) has the potential to improve uptake.\(^248\) Targeted campaigns, such as those being
implemented to boost Covid-19 vaccine uptake among specific ethnic minority groups in the UK, may also prove to be beneficial in reducing inequities in maternal vaccine uptake.

Prevention of bronchiolitis and chronic asthma

The majority of children become infected with RSV in early life, but only some infants develop bronchiolitis. In turn, only around 15% of children who are hospitalised with bronchiolitis have underlying conditions that are known to contribute to severity of infection (and an even smaller proportion of these are eligible for palivizumab prophylaxis against RSV infection). My third study concentrated in on one risk factor shown in the intermediary determinants section of Figure 8.1 (maternal smoking) to try to precisely estimate its contribution to inequities in admissions for bronchiolitis. I estimated that maternal smoking contributes to one fifth of the inequities in hospital admissions for bronchiolitis between infants. My fourth study was concerned with the relationship shown at the far right hand side of Figure 8.1. Again using a structured approach I precisely estimated the contribution of admission for bronchiolitis in the relationship between socioeconomic position and chronic childhood asthma. I show that targeting the prevention of bronchiolitis could reduce inequities in chronic asthma by almost one fifth. This further emphasises the importance of concentrating on interventions at the earliest point to prevent inequities in bronchiolitis and, in turn, childhood asthma.

Maternal smoking has long been identified as a risk factor for children’s respiratory ill health, and it continues to be on the public health policy agenda. Public Health England’s (PHE) strategy for 2020-25 specifically sets out specialist smoking cessation services for pregnant women to tackle social disadvantage in health. My findings add evidence to the importance of such policies that focus on maternal smoking (as well as other family members). On the other hand, my research also shows that 80% of the disparities in bronchiolitis admissions are not due to maternal smoking. More research is needed to understand the precise components of this 80%; however, reductions in exposure to air pollution and improving housing conditions in early life likely represent the most important avenues for early intervention. These risk factors are recognised in public health strategies for England and Scotland as contributing broadly to health inequities. PHE’s 2020-25 strategy specifically commits to lower air pollution to prevent health inequalities, and NHS Health Scotland’s 2017 strategy commits to improving suitable housing options. Evaluations of the implementation of these commitments and quantifying their impact on specific outcomes, including bronchiolitis and asthma, are an important next step.
Clinical interventions for bronchiolitis

My work also demonstrates that the previously observed increase in bronchiolitis admissions in England has continued through to 2016 (and likely beyond). In 2016, the rate of admission for bronchiolitis reached 58.4 per 1000 infant-years in England (95% CI 57.8 to 59.1). I show that these rates are even higher in Scotland (68.8 per 1000 infant-years in 2016, 95% CI 66.6 to 71.0). It has been suggested that hospital and healthcare policies such as changes in admission thresholds and accessibility of primary care services are an important driver of this increase, rather than changes in the transmissibility or severity of RSV. Hospital admission data are influenced not just by disease incidence and severity, but also by clinical decision-making, parental expectations and national healthcare policy (e.g. the availability of out of hours primary care services). Evaluations of alternative models of care for acutely ill infants, which reduce the need for hospital admission, could therefore also contribute to the reduction in bronchiolitis admissions. As argued by the British Lung Foundation (in reference to respiratory admissions on pediatric wards over winter) “much of this pressure could be eased by the development of decision support that would improve care across conditions”. The NHS Long Term outlines plans to address accident and emergency attendances in children using locally designed holistic services, which may have a beneficial effect on admission for bronchiolitis.

8.2.3. Future research

There are many avenues of further research to be exploited from the findings in this PhD. Many of these suggestions could be achieved through current research initiatives involving linkage between the administrative health data that I used in my thesis to cohort studies or other administrative and environmental datasets.

Geographical and socioeconomic variation in rates of bronchiolitis

Future investigations into the variation in rates of hospital admissions for bronchiolitis across England would profit from the inclusion of clinical factors unavailable in my datasets, such as a measure of admission thresholds. Including information on pathways to hospital admission would also be advantageous, particularly to understanding whether models of care for bronchiolitis differ by SEP. The incongruent patterns of admissions in London compared with the rest of the country is particularly warranted (and research on this is underway). Other important factors to consider in future investigations of geographical variation of RSV bronchiolitis epidemics include weather patterns and levels of common air pollutants. Studying the underlying pathogens that cause bronchiolitis will help understand whether the
socioeconomic gradient observed for all bronchiolitis admissions (mostly caused by RSV) are similar for other pathogens. This would require linkage to the Second Generation Surveillance System held by Public Health England or Electronic Communication of Surveillance in Scotland held by Public Health Scotland.\textsuperscript{474} This linkage is underway as part of the PICNIC study, which is a national birth cohort study on air pollution, housing and respiratory tract infections in children in England.\textsuperscript{475}

**Pathways between infant bronchiolitis and development of asthma**

I introduce statistical methods (and include the code for implementation) that can be used in future observational studies to investigate pathways between socioeconomic position and bronchiolitis and asthma. As causal inference methods advance and the potential to meaningfully investigate multiple pathways expands, teasing out the precise contribution of multiple mediators will become easier. Replication of the asthma/wheeze trajectory analyses over many study years is important for advancing understanding of trajectory-specific effects. Several identified confounders were not available in my studies, which ought to be included in future research on the association between bronchiolitis and asthma are outdoor air pollution, housing conditions (including damp and mould, overcrowding and indoor air quality) and breastfeeding.\textsuperscript{196,199,476} These risk factors should additionally be investigated as mediators in the relationship between SEP and asthma to further inform the focus and timing of interventions. This is a particular priority for air pollution, a risk factor with wide public awareness and an ever growing evidence base,\textsuperscript{149,477} but for which the precise early-life contribution to long-term disease risk (particularly asthma) remains to be quantified.\textsuperscript{478}

### 8.3. The impact of the SARS-CoV-2 pandemic

At the time of writing my discussion, the SARS-CoV-2 pandemic is ongoing. The hospital admission and death rate due to this virus and the resulting disease, Covid-19, are particularly high in the UK relative to most other countries around the world. As at 1 March 2021, the UK has the fifth highest recorded number of Covid-19 deaths of any country in the world.\textsuperscript{479} The effects of the pandemic on public health more broadly in the UK will take some time to fully emerge; however, there are some immediate factors that ought to be acknowledged in relation to the findings in my thesis.

At the same time as SARS-CoV-2 continues to circulate, albeit at lower levels than in the autumn and winter of 2020, other respiratory viruses had far lower circulation compared to other years. Surveillance information collated through the Respiratory Datamart system and
published by Public Health England, showed that RSV positivity was 0.0% in week 50 (2020) compared to 13.5% in the same week a year before. Research published from Western Australia shows a 98% decrease in RSV detections in children through winter 2020. Recent Hospital Episode Statistics for Admitted Patient Care and Outpatient Data (HES APC) statistics published by NHS Digital do not include information on diagnoses, but point to the implications on hospital admissions for bronchiolitis. This provision data show a 55.9% decrease in emergency hospital admissions among 0-4 year olds in December 2020 ($N = 61,707$) compared to December 2019 ($N = 27,234$).

These changes in admission patterns for respiratory viruses in children strongly highlight the utility of simple behavioural measures in preventing infectious disease, and has the promise of changes to the transmission of viruses in the future (if sustained). Hand washing is a prominent example, which is a very effective measure to prevent respiratory infection in children. As it is recognised by paediatricians that bronchiolitis places a major burden on NHS paediatric wards and intensive care units between October and March, a larger group of at risk children were recommended to receive palivizumab prophylaxis in 2020/21 to keep them out of hospital. However, these patterns of RSV also have the potential to have negative repercussions in future years. It is hypothesised that, partially due to a larger proportion of immunologically naïve children, the next RSV epidemic may be larger than previous years, leading to greater morbidity from infection. Prominently, this period of sudden change offers a natural experiment to examine whether early bronchiolitis actually leads to asthma.

### 8.4. Concluding remarks

The stark socioeconomic gradient in the incidence of bronchiolitis and the prevalence of asthma are major concerns for child public health in the UK. Intervening on maternal smoking during pregnancy continues to offer an important avenue for intervention, particularly to reduce disparities in infant bronchiolitis infection. I have also demonstrated, using novel application of causal inference methodology, that preventing severe bronchiolitis could reduce the socioeconomic gradient in chronic asthma prevalence by nearly one fifth. This thesis offers a guide to the use of causal inference methods in the study of inequities in health using complex, linked administrative health data. Future work including indicators of air pollution and housing conditions is particularly warranted in the complex unravelling of pathways to bronchiolitis and subsequent childhood asthma.
APPENDIX 0: RESEARCH OUTPUTS

Appendix 0.1. Publications during PhD period

Research papers


Peer reviewer

November 2020 Scientific Reports

May 2019 The Journal of Epidemiology and Community Health

May 2019 BMC Pulmonary Medicine
Appendix 0.2. Conferences and presentations

12-13 September 2019  UCL-Birkbeck MRC DTP retreat. Poster entitled ‘Defining asthma/wheeze groups using symptom trajectories’


15-19 October 2018  UK China Workshop on Childhood Respiratory Disease, Chongqing (China). Poster presentation entitled ‘Seasonal and geographical variation of severe bronchiolitis across England’. Funded via the Newton Fund.

Appendix 0.3. Secondment

January to July 2020  Fact checker on the editorial team, Full Fact

During this 6-month secondment at the UK’s independent, non-partisan, fact checking charity, I researched and published articles on the misreporting of health research in the mainstream media, on social media and by public figures. The majority of these focussed on misinformation about Covid-19 and included several longer reads on topics related to my PhD. For example, how does the new coronavirus affect pregnancy? and what do we know about Covid-19 inequalities among people from minority ethnic groups?
Appendix 1.1. EJE publication: Mother’s education and offspring asthma risk in 10 European cohort studies

Mother’s education and offspring asthma risk in 10 European cohort studies

Kate Marie Lewis1 · Milagros Ruiz1 · Peter Goldblatt2 · Joana Morrison1 · Daniela Porta1 · Francesco Forastiere1 · Daniel Hryhorczuk4 · Oleksandr Zvinchuk5 · Marie-Joseph Saurel-Cubizolles6 · Sandrine Lioret7 · Isabella Annesi-Maesano8 · Martine Vrijheid1,9,10,11 · Matias Torrent1,9,11,13 · Carmen Iniguez9 · Isabel Larrauriag1,4,15 · Margreet W. Harskamp-van Ginckel16 · Tanja G. M. Vrijkot16 · Jana Klanova1,17 · Jan Svancara18 · Henrique Barroso19,20 · Sofia Correia19,20 · Marijo-Klita Jarven15 · Anja Tanulich21,22,23,24 · Johnny Ludvigsson21 · Tomas Farøj22 · Michael Marnot12 · Hynck Pihart3

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Abstract Highly prevalent and typically beginning in childhood, asthma is a burdensome disease, yet the risk factors for this condition are not clarified. To enhance understanding, this study assessed the cohort-specific and pooled risk of maternal education on asthma in children aged 3–8 across 10 European countries. Data on 47,099 children were obtained from prospective birth cohort studies across 10 European countries. We calculated cohort-specific prevalence difference in asthma outcomes using the relative index of inequality (RII) and slope index of inequality (SII). Results from all countries were pooled using random-effects meta-analysis procedures to obtain mean RII and SII scores at the European level. Final models were adjusted for child sex, smoking during
pregnancy, parity, mother’s age and ethnicity. The higher the score the greater the magnitude of relative (RII, reference 1) and absolute (SII, reference 0) inequity. The pooled RII estimate for asthma risk across all cohorts was 1.46 (95% CI 1.26, 1.71) and the pooled SII estimate was 1.90 (95% CI 0.26, 3.54). Of the countries examined, France, the United Kingdom and the Netherlands had the highest prevalence’s of childhood asthma and the largest inequity in asthma risk. Smaller inverse associations were noted for all other countries except Italy, which presented contradictory scores, but with small effect sizes. Tests for heterogeneity yielded significant results for SII scores. Overall, offspring of mothers with a low level of education had an increased relative and absolute risk of asthma compared to offspring of high-educated mothers.

Keywords Asthma · Children · Cohort studies · Maternal education · Socioeconomic position · Disease risk

Introduction

Asthma is a chronic disease of the bronchial tubes in the lungs, which typically develops in childhood, and is characterised by wheezing, breathlessness, chest tightness and coughing [1–3]. Children with this condition face increased absence from school, reduced family life participation and lower quality of life compared to non-asthmatic peers [4]. Asthma is one of the most common diseases amongst children worldwide [1] and is particularly prevalent in Western Europe (9.7% asthma ever in 6–7 year olds) [5]. With the appropriate measures in place most cases of asthma can be controlled, but no proven prevention or cure currently exist [1, 6]. With financial costs for asthma in EU countries estimated at €72.2 billion per annum, the case for improved understanding is strong [7].

The causes of childhood asthma are multifaceted, with genetic and environmental exposures increasing susceptibility to the development of this condition [5, 8]. Known early life risk factors include preterm birth, infant weight gain and adiposity, in utero and postnatal tobacco smoke exposure, household damp or mould, lower respiratory infections and pollutants such as NOx, sulphur dioxide and particulate matter [8–12]. As socioeconomically deprived groups face increased exposure to these health damaging risk factors, grasping how social determinants impact on asthma represents a broader framework for establishing preventative measures. Prospective cohort studies in Sweden, the UK and the Netherlands have previously linked parental income, occupation and education to the increased development and severity of childhood asthma [13–17]; however, this evidence is restricted to a handful of European countries and has limited comparability due, in part, to the use of inconsistent socioeconomic position indicators [18].

This research contributes to current understanding through analysis of maternal education, arguably the most important social determinant of early child health [19], as a risk factor for early childhood asthma across Europe. Europe is highly diverse, with wide variation in social, economic and health policies and outcomes between and within countries [20, 21]. Comparing differences in asthma inequalities using a common indicator across the continent will help to decipher contextual factors and enable further consideration of the pathways to asthma development. Early childhood is a crucial developmental period of life and given that asthma trajectories can be set in place as early as 6/7 years old, an optimal time for intervention [17, 18]. We hypothesised that social inequalities in asthma risk would differ between European countries, but broadly follow an inverse relationship with mother’s education level.

Methods

Data sources

Drawing on research by the DRivers for Health Equity research programme [22], the current study used data from 10 European birth cohort studies (Box 1). In total, complete data was available for 47,099 singleton children born between June 1985 and January 2008. The sample comprised of cohorts (in descending order of size) from: England and Wales (UK-MCS, n = 13,829); Southeast Sweden (SE-ABIS, n = 8,308); Oulu and Lapland, Finland.
### Appendix: Chapter 1

**Box 1 Cohort study locations, names and cohort members’ year of birth**

<table>
<thead>
<tr>
<th>Location</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>FI-NFBC</td>
<td>Finland—Northern Finland birth cohort 1985-1986 study</td>
</tr>
<tr>
<td>FR-EDEN</td>
<td>France—the mother–child study of pre- and postnatal determinants of child growth, development, and health, 2003-2006</td>
</tr>
<tr>
<td>IT-GASPII</td>
<td>Italy—the gene and environment prospective study on infancy in Italy, 2003–2004</td>
</tr>
<tr>
<td>NL-ABCD</td>
<td>The Netherlands-Amsterdam born children and their development study, 2003–2004</td>
</tr>
<tr>
<td>PT-G21</td>
<td>Portugal—the generation XXI study, 2005–2006</td>
</tr>
<tr>
<td>ES-INMA</td>
<td>Spain—the environment and childhood project, 1997–2008</td>
</tr>
<tr>
<td>SE-ABIS</td>
<td>Sweden—all babies in Southeast Sweden, 1997–1999</td>
</tr>
<tr>
<td>UA-FCOU</td>
<td>Ukraine—the family and children of Ukraine study, 1992–1996</td>
</tr>
<tr>
<td>UK-MCS</td>
<td>United Kingdom—Millennium cohort study, 2000–2001</td>
</tr>
</tbody>
</table>

(FI-NFBC, n = 6892); Porto, Portugal (PT-G21, n = 6486); Amsterdam, the Netherlands (NL-ABCD, n = 3387); Brno and Znojmo, Czech Republic (CZ-CELSpac, n = 2801); Gipuzkoa, Menorca, Sabadell and Valenciac, Spain (ES-INMA, n = 1843); Kyiv, Dneprodzerzhinsk (now Kaminksi) and Mariupol, Ukraine (UA-FCOU, n = 1757); Nancy and Poitiers, France (FR-EDEN, n = 1159); and Rome, Italy (IT-GASPII, n = 637).

Ethical approval and participant consent had been approved by each cohort study before data collection, and no further ethical approval was required for the current study. Anonymisation and extraction of the relevant data was performed by researchers from each cohort study before combining with other datasets for analyses. For further cohort details, see previous research [22].

### Study variables

The definition of childhood asthma varied across the 10 datasets. SE-ABIS retrieved doctor diagnoses directly from primary care and hospital records at age 5 (ICD-10 code J45). All other studies determined asthma presence (yes/no) by mothers’ answer to a survey question, which asked whether their child had had a doctor diagnosis of asthma (FR-EDEN age 5, UA-FCOU age 7, PT-G21 age 4/5); asthma as a ‘long term illness’ (FI-NFBC age 7/8) or an ‘allergic disease’ (CZ-CELSpac age 3); ever had asthma (NL-ABCD age 7/8, IT-GASPII age 7, UK-MCS age 5); or had asthma in the last 12 months (ES-INMA age 3/4). Each cohort study ascertained all other information relevant to the current analyses through mother’s self-report prior to their child’s birth, shortly after birth or when the child was 9-months old (UK-MCS only).

Maternal education level was determined with a variety of questions, including years (FI-NFBC; FR-EDEN; PT-G21; NL-ABCD) and stage (ES-INMA; IT-GASPII; SE-ABIS; UA-FCOU) of schooling completed and highest academic qualification (UK-MCS). CZ-CELSpac and UA-FCOU enquired about highest education level out of the parents and grandparents. Education levels across countries were converted to the same scale using the international Standard Classification of Education (ISCED 97) [23]. We banded the seven levels of the ISCED 97 into three: low, up to lower secondary education (ISCED 0-2); medium, upper secondary education (ISCED 3); and high, post-secondary education and higher (ISCED 4-6).

Several potential confounding factors were present across all cohorts and were therefore included as covariates in the final models. Maternal age at child’s birth (grouped into <30/30 years old), sex of the child (male/female), parity, which we dichotomised into yes (nulliparous) or no (multiparous) by mother’s report of whether the current birth was their first live-born child; Maternal ethnicity, dichotomised into prevalent ethnicity of the country/region—defined as Caucasian White (FR-EDEN; IT-GASPII), White (UK-MCS), European (CZ-CELSpac) or by country of birth (NL-ABCD; ES-INMA) or minority ethnicity for all women not meeting the definition. Smoking during pregnancy (yes/no) based on the following definitions: ≥one cigarette a day during pregnancy (ES-INMA); smoking throughout pregnancy (UK-MCS); smoking regularly in last 6 (CZ-CELSpac) or 12 months (UA-FCOU); smoked in 1st or 3rd trimester (PT-G21); smoked at the beginning of pregnancy (FI-NFBC); or smoked at all during pregnancy (IT-GASPII; FR-EDEN; NL-ABCD; SE-ABIS). Non-singleton children (n = 1399) were excluded from further analyses.

### Data analysis

All analyses were conducted in Stata 14 [24]. Missing variables data ranged from 3.2% (sex) to 27.7% (asthma) and missingness in the outcome variable was significant related to lower education level and smoking in pregnancy (p < 0.001). To enable comparison across cohorts, cases without complete data for all study variables were excluded.
from analyses. Sampling and attrition survey weights, available for UK-MCS, and generic weights for non-MCS cohorts (probability weight = 1) were applied to all analyses [18]. The distribution of mothers’ education for each cohort was age-standardised using the WHO European Standard Population [25].

Prevalence of asthma by mother’s education was calculated for each cohort. The \( \chi^2 \) test for trend assessed linearity across educational groups with the exception of UK-MCS, to which linear regression was applied to survey weighted data (and compared to unweighted \( \chi^2 \) test results). The Relative and Slope Indices of Inequality (RII/ SII) were used to compute cohort-specific and total differences in asthma outcome [22, 26, 27]. RII represents the relative change in asthma outcome between highest and lowest education group (the prevalence ratio), whilst the SII is the absolute change (the prevalence difference). A score higher than 1 (RII) and 0 (SII) indicates inequity, and the higher the score the greater the magnitude of the inequity.

RII and SII scores, adjusted for child sex, first birth, mother’s age at birth, mother’s ethnicity and maternal smoking status during pregnancy were determined for each cohort. We pooled the results from each cohort using meta-analysis procedures to obtain mean RII and SII scores at the European level. Assuming between-cohort heterogeneity, we applied a random effects model. The \( I^2 \) measure was used to test the extent to which heterogeneity was present. Lastly, we conducted sensitivity analysis comparing the results of fixed and random effects upon the models.

### Results

Table 1 presents characteristics of the final sample. Of the 47,099 children, 47.9% were female, 50.9% of mothers were younger than 30 at their child’s birth and 46.3% was their first live birth. UK-MCS and ES-INMA had the highest rates of smoking in pregnancy, 33.0 and 29.7% respectively, compared to 7.8% in NL-ABCD and 8.8% in SE-ABIS. The proportion of mothers from ethnic minorities ranged from 0% (FI-NFBC; SE-ABIS; UA-FCOU) to 14.5% (NL-ABCD). The gap between proportion of mothers in low and high ISCED education levels was largest in IT-GASPII (13.2 vs. 36.0%) and PT-G21 (47.2 vs. 25.4%). Asthma was identified in 37.7% (79.9%) children with the prevalence ranging from 1.7% (UA-FCOU) to 14.4% (UK-MCS) across cohorts. As shown in Table 2, the unadjusted trend of increased asthma prevalence with lower maternal education level was strongest in UK-MCS (\( p < 0.001 \), NL-ABCD (\( p < 0.001 \)) and FR-EDEN (\( p = 0.008 \)) cohorts. A weaker trend was noted for SE-ABIS (\( p = 0.031 \)). IT-GASPII showed a reverse pattern, with asthma more prevalent in the highest education level, however, this results showed no significant trend (\( p = 0.60 \)). There was little to no evidence of an educational trend across all other cohorts, which, in some cases, may be due to a lack of statistical power to detect an effect.

As displayed in Fig. 1a, the inverse relative risk of asthma with maternal education is strongest in cohorts FR-EDEN (RII 2.07 95% CI 1.10, 3.89), UK-MCS (RII 1.73 95% CI 1.44, 2.09) and NL-ABCD (RII 1.65 95% CI 1.00, 2.72). This pattern is reflected in absolute inequity values (Fig. 1b); Children in the FR-EDEN cohort have a 9.0% risk difference of asthma, UK-MCS a 7.3% difference and NL-ABCD a 5.9% difference. CZ.CELEPAC, FI-NFBC, ES-INMA, PT-G21, SE-ABIS and UA-FCOU have

| Table 1 Characteristics of study sample, overall and cohort-specific |
|-----------------------------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|
|                             | Total           | CZ              | FI              | FR              | IT              | NL              | PT              | ES              | SE              | UA              | UK              |
| Sample size (N)             | 47,099          | 2801            | 6892            | 1159            | 637             | 3387            | 1843            | 6486            | 8308            | 1757            | 13,829          |
| Asthma: yes (%)             | 7.9             | 1.9             | 5.2             | 11.7            | 8.2             | 9.1             | 2.7             | 4.3             | 6.7             | 1.7             | 14.4            |
| Child’s sex: female (%)     | 47.9            | 47.0            | 49.1            | 47.0            | 49.1            | 48.9            | 48.6            | 49.2            | 48.3            | 47.0            | 48.9            |
| Mother’s ethnicity: minority (%) | 4.3             | 0.7             | 0.0             | 2.1             | 2.0             | 2.0             | 1.45            | 4.4             | 4.2             | 5.0             | 8.9             |
| Mother’s age at birth: <30 (%) | 50.9            | 86.4            | 66.0            | 46.4            | 20.7            | 24.5            | 35.3            | 50.4            | 43.7            | 84.3            | 48.6            |
| Firstborn: yes (%)          | 46.3            | 48.2            | 34.3            | 46.4            | 58.6            | 58.0            | 54.8            | 57.1            | 40.2            | 67.2            | 43.2            |
| Maternal smoking: yes (%)   | 21.0            | 20.4            | 18.5            | 19.3            | 11.0            | 7.8             | 29.7            | 22.5            | 8.8             | 10.8            | 33.0            |
| Mother’s education          |                 |                 |                 |                 |                 |                 |                 |                 |                 |                 |                 |
| Low (%)                     | 24.1            | 35.6            | 25.2            | 4.2             | 13.2            | 8.3             | 29.7            | 47.2            | 33.7            | 5.2             | 12.2            |
| Medium (%)                  | 43.5            | 30.1            | 51.2            | 34.9            | 50.9            | 20.4            | 38.9            | 27.5            | 32.1            | 78.4            | 57.3            |
| High (%)                    | 32.4            | 25.3            | 23.7            | 60.9            | 36.0            | 71.3            | 31.4            | 25.3            | 34.2            | 16.3            | 30.5            |

*Weighted proportions
positive RII and SII values, but with smaller statistically insignificant effect sizes. IT-GASHI displays a contrary pattern; low education level is associated with 34% lower odds (RII = 0.66 95% CI 0.24, 1.33) and 4.68% decreased risk difference (SII = -4.68 95% CI -12.92, 3.56) of asthma compared to high education. However, these relationships are not statistically significant.

Across all European cohorts, children born to mothers in the lowest tertile of education were 46% more likely to develop asthma than children born to mothers in the highest tertile (95% CI 1.26, 1.71). The pooled estimate of SII was 1.90 (95% CI 0.26, 3.54), indicating a 1.9% mean risk difference of asthma between children born to mothers in the low compared to high educational group. An $I^2$ value of 9.2% ($p = 0.36$) in the RII model gives evidence of low heterogeneity in relative scores between cohorts; however, absolute scores (SII) were moderately to highly heterogeneous ($I^2 = 72.0\%, p < 0.0001$) [28]. Fixed effects meta-analysis techniques slightly increased relative estimates (RII = 1.50 95% CI 1.32, 1.71), whilst decreasing absolute estimates (SII = 1.47, 95% CI 0.70, 2.34).

**Discussion**

This study uniquely combined prospective data of educational inequalities in childhood asthma from 10 European cohort studies. Overall, we found relative and absolute asthma risk inversely related to maternal education, which is broadly consistent with previous research [8, 14, 29–32]. However, the magnitude of difference varied across countries and substantial heterogeneity was present in absolute differences. The highest prevalence of asthma and most pronounced inequity were observed in Western European countries.

France, the UK and the Netherlands cohorts had asthma rates between 9 and 14%, compared to less than 2% of the Czech Republic and Ukraine cohorts. East-to-West geographic variations have been noted previously, although are now reportedly diminishing [1]. Whilst climatic risk factors such as low altitude, temperature variations and relative humidity may explain some of the difference, it is argued that the urban ‘western lifestyle’ is a major contributor—and the adoption of this by Eastern Europe the reason for the closing gap [33, 34]. Western lifestyle factors include increased exposure to vehicle emissions and a diet high in calories, fat and sugar, but low in fruit and vegetables.

Maternal education affects child health through several mechanisms. Lower levels of education are linked to lower employment positions, fewer material resources and social support, as well as limited access to health information (and a diminished ability to act on this information) [19, 35, 36]. Prominently, this leads to greater exposure to health damaging risk factors for the mother and consequently their child. Although our analyses were adjusted for several potential risk factors, data constraints meant that many other risk factors were not included. It is likely that a large proportion of inequity may be explained by differences in exposure to risk factors such as tobacco smoke after birth, infant weight gain and adiposity, household damp, infections, and pollution [8–12]. Family history of asthma and atopy, unmeasured in this study, is one of the strongest risk factors for childhood asthma [8]. However, this does not fully explain why the impact of
Fig. 1: Forest plots of the RII (a) and SII (b) in childhood asthma across 10 European cohorts (squares) and combined (diamond), adjusted for child sex, smoking during pregnancy, parity, mother’s age and ethnicity; values to the right of the undashed vertical line indicate greater inequity for low compared to high maternal education groups.
maternal education is greater in some countries compared to others.

Explaining difference in inequity across countries requires us to look at the structural facets that constrain and disadvantage people with lower education [37]. It is proposed that the intergenerational impact of education is additionally conflated by the broader socio-political and cultural landscape within countries [38]. The results from Western Europe suggest that whilst access to health care and social security is important, it is not enough to counter inequities [38]. Universal provision in healthcare, in particular, does not mean universal uptake [29]. For example, disadvantaged groups in the UK are less likely to avail of health promotion and prevention services [39]. Anxo et al.’s [40] review found that the impact of women’s educational attainment on employment integration is more pronounced in France, Italy and Spain compared to Sweden [40]. Higher employment rates and macro-social policies that value women and families in the Nordic countries may partially explain lower levels of inequity compared to Western Europe [41, 42]. However, high rates of adiposity amongst children of parents with low education persist in Sweden and Finland despite egalitarian policies [37, 43].

The Mediterranean diet may protect against the impact of maternal education in Southern Europe [44].

Birth cohort studies are an essential and unique source of information on the multifaceted predictors of chronic disease [47]. Combining 10 European studies has enabled a comparison of more than 47,000 children over several years of life. The prospective design addresses issues of reverse causation inherent to cross-sectional studies. In addition, as the results were not reliant on published data, this study avoided publication bias. However, several limitations must be acknowledged when considering the current study. Firstly, rates of asthma and the inequities shown are not necessarily indicative of the country or area in which each cohort study were based. Many cohorts sampled only one major urban area and we may expect regional differences in rates of education in rural areas, for instance.

Combining multiple datasets brings difficulties and, as highlighted by the heterogeneous absolute rates of inequity, there is an unexplained difference across the included cohorts. Unmeasured confounding, such as the risk factors mentioned above, is likely to contribute to this. In addition, several study aspects were not consistent across cohorts. Age at which asthma was defined ranged from 3 to 8 years old, and the timespan in which to consider a diagnosis from the last 12 months to at any point across the child’s lifetime. Symptomatology of asthma, as well as the impact of inequity on the disease outcome, can differ across age in childhood [1, 14, 46]. Birth year ranged from 1985 to 2008 across the cohorts, adding time-varying contextual factors to the findings. For example, the Spanish cohort study recruited children over a decade, encompassing a period of economic growth and better perinatal outcomes than in the period of economic recession that followed [19].

The proportion of mothers with low education was inconsistent across studies. Both low education and young mothers appear underrepresented in the Italian sample, which may imply limited study power and explain the contradictory results shown by this cohort. Previous cross-sectional research in Rome found a significant trend between lower paternal education and increased risk of asthma, and a similar but insignificant trend of maternal education [47]. The proportion of mothers with low education was also low in France, Ukraine and the Netherlands; however, these are similar to national averages for the time [22]. Highest education level in the Ukraine and Czech Republic cohorts were based on several family members, which likely conflated the education level of some mothers. Almost all cohorts relied on maternal reports of asthma rather than doctor’s diagnosis, which likely led to imprecise estimations of the condition. Cesaroni et al. [47] speculate that parents from higher socio-economic positions are more likely to report asthma especially where symptoms are mild. Missing data for the asthma outcome was significantly related to lower maternal education across the dataset, further indicating a socially patterned bias to the results.

Despite limitations, this comparative analysis across European cohorts demonstrates that maternal education is implicated in offspring asthma risk. The difference in magnitude of inequity across countries, suggests contextual factors affect the translation between maternal education and offspring health outcomes and, importantly, that reduction in inequity is possible. Given that early disadvantage in respiratory health puts into place a negative trajectory that can be difficult to modify [17], these results reinforce the need for early, if not pre-birth, interventions to offset the burden of this disease [18, 38]. Universal and high-quality perinatal care including information about asthma risk that is clearly and respectively articulated is a necessity [48, 49]. Both improving the educational attainment of young women and combating systematic barriers faced by women with lower education, such as access to employment, is essential to ensure that these offspring are not further disadvantaged [40, 42].

Future cross-European analyses could be improved with the inclusion of known environmental risk factors for asthma and by the use of standardised measures (such as the ISAAC for asthma) to increase comparability and enable pathway analyses [5, 17]. Exclusion of genetic predispositions using genetic methods, such as twin and family studies, will clarify environmental causes with more precision. Differentiating between asthma phenotypes and
additional indicators of socioeconomic position, such as income, will help further untangle cross-country differences [29, 50].

Acknowledgements All phases of this study were supported by a European Union’s Seventh Framework Programme grant, 27850, as part of The Determinants to Reduce Health Inequity Vis Early Childhood, Realising Fair Employment, and Social Protection (DRIVERS) research programme. The Czech ELSPAC Study (CZ-ELSPAC) was supported by the Ministry of Education of the Czech Republic: CETOCOPEN plus project (CZ.02.1.01/0002/15/000 000469) and RECETEX Research Infrastructure (LM2015051). The Northern Finland Birth Cohort (FNBC5586) received financial support from the Academy of Finland; Biocenter, University of Oulu, Finland; the European Commission EUROBCS, Framework 5 Award (QLGI-CT-2000-01643); EU P97 EurHEALTH4aging- 277149; the Medical Research Council, UK (BveMetsSynSAVLVE); and the NRC Centenary Early Career Award. The Amsterdam Born Children and their Development Study (NLABCD) received funding from the Netherlands Organization for Health Research and Development (ZonMW) Grant (TOP 40-008.12-98-11010). The All Babies in Southeast Sweden Study (SE-ABS) has received financial support from the Juvenile Diabetes Research Foundation, Swedish Child Diabetes Foundation (Ram diabetesfonden). The Research Council of South-nor Sweden (FORSS), Swedish Research Council K2005-72X- 11242-11A, and Al-Fat County Council of Östergötland. The INMA study was funded in part by grants from the European Union (FP7- ENV-2011-28257 and HEALTH2010.2.4.5.1), Spain (Instituto de Salud Carlos III and The Ministry of Health), the Conselleria de Sanitat of the Generalitat Valenciana, Department of Health of the Basque Government, the Provincial Government of Galicia, and the Generalitat de Catalunya-CIRIT. Family and Children of Ukraine (UA-FCQ) study was supported by US NIH Fogarty International Center and National Academy of Medical Sciences of Ukraine. KML is funded by a Medical Research Council UK doctoral training fellowship.

Compliance with ethical standards

Conflict of interests The authors declare that they have no conflict of interests.

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References

Appendix 3.1. Linking Mother and Infant Data in HES APC

The Hospital Episode Statistics Admitted Patient Care (HES) dataset contains routinely collected maternity records from NHS hospitals across England.\textsuperscript{45} There are two types of HES maternity records available, the (maternal) delivery record and the (baby) birth record.\textsuperscript{273} Maternal and baby records in HES do not share a unique identifier, however, and are not linked together in the dataset we receive from NHS Digital. Linking delivery and birth records leads to a more complete medical record for the mother and child, improving reliability and statistical power of secondary research using this data. To connect mother and baby records for HES years 2013/14 to 2016/17, I used deterministic and probabilistic linkage methods as outlined by Harron et al.\textsuperscript{273}

Deterministic linkage is relatively straightforward. It involves applying a set of rules to classify pairs, in this case mothers and infants, as links or non-links.\textsuperscript{485} For this dataset, GP practice, infant sex, birthweight, gestational age, mother’s age and birth order were used to deterministically match pairs (see Table A3.1 for more details). Improvement in hospital recording of data means that rates of deterministic linkage between mothers and babies have grown over the years;\textsuperscript{45} however, in 2016/17 this type of linkage still remained below 50%. For the remaining mother and infant records, probabilistic linkage was applied, where the likelihood of records belonging together is represented by a match weight. This method is suitable for matching records that contain several non-unique identifiers, which may, in combination, uniquely identify an individual.\textsuperscript{273} These variables are referred to as ‘quasi-identifiers’\textsuperscript{486} and are listed for the current dataset in Table A3.1.

Steps for probabilistic linkage

Two probabilities are needed to calculate match weights:

- **U-probabilities** - the likelihood that (non-missing) variables across two records randomly match.
- **M-probabilities** are the probability there is an agreement on a (non-missing) variable given that two records are a true match.

I calculated U-probabilities by selecting a random sample of 5000 unlinked mother and baby records within each HES year, and running pairwise comparisons of each variables across all these records (i.e. 25,000,000 comparisons).\textsuperscript{273} M-probabilities were calculated using the
deterministically linked records (as outlined above). Weights were then calculated for each variable across record pairs using the formula, \( \log_2(M\text{-probability}/U\text{-probability}) \), and summed to get an overall match weight for each pair.

Match weights across record pairs were reviewed using plots and manual review to select a single threshold. All pairs above this values were deemed to be true matches and these were placed into the deterministically linked dataset. The process of calculating M-probabilities and match weights was repeated on the new dataset. I carried out three repetitions of these steps in total to ensure match weights had stabilised. The final iteration involved applying the now stabilised M-probabilities to all non-deterministically linked pairs to get the final match weights. Pairs below the final threshold were placed into a separate dataset. Pairs above the threshold were ordered by descending match weight and the highest-weighted mother for each baby was selected, followed by the highest weighted baby for each delivery record (mothers may have more than one singleton baby in a year).

**Table A3.1.** Details of variables used in linkage

<table>
<thead>
<tr>
<th>Variable</th>
<th>HES name</th>
<th>Description and linkage type (D = Deterministic and P = probabilistic)</th>
</tr>
</thead>
<tbody>
<tr>
<td>GP practice</td>
<td>gpprac</td>
<td>The code of the general practice where the patient’s registered GP works</td>
</tr>
<tr>
<td>Provider code</td>
<td>procode3</td>
<td>The first 3 characters of the organisation acting as the health care provider</td>
</tr>
<tr>
<td>Infant sex</td>
<td>sexbaby</td>
<td>Sex of the baby</td>
</tr>
<tr>
<td>Birth order</td>
<td>birordr</td>
<td>Position in sequence of birth (i.e must be 1 for singleton births)</td>
</tr>
<tr>
<td>Birth weight</td>
<td>birweit</td>
<td>Weight of the baby in grams immediately after birth</td>
</tr>
<tr>
<td>Gestational age</td>
<td>gestat</td>
<td>Length of gestation in weeks</td>
</tr>
<tr>
<td>Mother’s age</td>
<td>matage</td>
<td>Mother’s age at delivery</td>
</tr>
<tr>
<td>Delivery place</td>
<td>delplac</td>
<td>Delivery place [2013/14 only]</td>
</tr>
<tr>
<td>Postcode district</td>
<td>postdist</td>
<td>First 3/4 characters of postcode [2014/15 onwards]</td>
</tr>
<tr>
<td>Postcode area</td>
<td>postdist</td>
<td>First character of postcode</td>
</tr>
<tr>
<td>Gestation period</td>
<td>anagast</td>
<td>Gestation period in weeks at first antenatal assessment</td>
</tr>
<tr>
<td>Delivery place (intended)</td>
<td>delinten</td>
<td>The intended place of delivery</td>
</tr>
<tr>
<td>Delivery method</td>
<td>delmeth</td>
<td>defines the method used to deliver a baby that is a registrable birth</td>
</tr>
<tr>
<td><strong>Delivery method</strong> (onset)</td>
<td>delonset</td>
<td>Defines the method used to induce (initiate) labour</td>
</tr>
<tr>
<td>-----------------------------</td>
<td>----------</td>
<td>----------------------------------------------------</td>
</tr>
<tr>
<td><strong>Anaesthetic</strong> (post)</td>
<td>delposan</td>
<td>Anaesthetic given post labour or delivery</td>
</tr>
<tr>
<td><strong>Anaesthetic</strong> (during)</td>
<td>delprean</td>
<td>Anaesthetic given during labour or delivery</td>
</tr>
<tr>
<td><strong>Delivery status</strong></td>
<td>delstat</td>
<td>Status of person conducting the delivery (e.g. hospital doctor, midwife)</td>
</tr>
<tr>
<td><strong>Resuscitation method</strong></td>
<td>biresus</td>
<td>Method to get the baby breathing</td>
</tr>
<tr>
<td><strong>Birth status</strong></td>
<td>birstat</td>
<td>Live or still birth</td>
</tr>
<tr>
<td><strong>Ethnicity</strong></td>
<td>ethnos</td>
<td>Ethnicity of the patient</td>
</tr>
</tbody>
</table>
Appendix 3.2. Data linkage feedback form

This form was sent to the Great Ormond Street hospital parent carer advisory group and advertised through Asthma UKs monthly bulletin for parents and carers.

10th November 2017

Using data to understand risk factors for asthma in children

Who am I and what is my research?

I am a research (PhD) student from UCL Great Ormond Street Institute of Child Health.

My research project is focussed on understanding the early risk factors leading to different types of childhood asthma.

To do my research I will use electronic health data from hospitals in England and Scotland.

Using data on the whole childhood population in these countries means that I get a more complete picture of hospital admissions for asthma and the early risk factors that lead to this. For example, early respiratory infection and premature birth.

What do I want to do?

To identify more children with asthma, we want to link hospital admission data to drug dispensing data from pharmacies. I am interested in asthma medication in particular. This will improve my research by including more children with different types and severities of asthma. FIGURE 1, below, shows the linked data required for my research project. This type of linkage is only available for children in Scotland at the moment.
3.0 How can you help?

I would like you to consider this research project and the data linkage proposed above. The process of secure data linkage is explained in FIGURE 2, on the next page. Then I would like your feedback, by answering the question below.

**Question for you:** Do you have any thoughts or concerns on combining these sources of data for this project and if so why?

Any other questions for me?

If you would like to find out more about this research, email [email]...[email]

Thank you for taking the time to answer these questions, your help is invaluable to my work.
FIGURE 2: Process of secure data linkage

What about consent?

We do not receive information that can directly identify a patient, like names, postcode or NHS number. We require special permissions from the NHS and the Office for National Statistics to allow these data to be linked. We can only use the data to answer specific research questions. We are using the data without specific consent.

Steps to data linkage:

1. Personal information sent to linkage server
2. Personal information linked and returned
3. Linked data, not personal information, sent to the researcher (me!)
Appendix 3.3. Feedback request sent to the Farr Institute Public Panel

The following email was sent to members of the Farr Institute Public Panel alongside an earlier draft of the patient opt out commentary article presented in Appendix C. We received 10 responses from the panel, which helped shape the points within the article.

__________________________________________________________________________________

4th July 2018

I am a PhD student looking into the impact of deprivation on childhood respiratory illness. To do this work I use hospital data from babies (linked to their mother’s data) across England. This provides me with information on various risk factors for respiratory illness including preterm birth, deprivation, area of the country and early childhood illness. The linked maternal data is very important as it enables me to look at a wider range of risk factors than baby data alone would provide, such as rates of pollution in the local area of residence and delivery type. It is very important to have complete whole population data in this work so that I can understand the full picture and look at differences across the country. However, in the newest years of hospital data there are more and more instances where there is no maternal information available at all, which means that critical information (such as pollution and deprivation) are not available for some babies.

After speaking with colleagues, we realised that the increase in missing data is largely due to patients choosing to ‘opt-out’ of sharing their data for purposes other than clinical care (such as for research). Research on this topic shows that there is little information provided for the public to understand how and why their data may be used, and the safeguards in place to protect their information. This finding led me to write an article to explain these problems, with specific reference to the implications for maternal statistics. The British Medical Journal have expressed interest in this topic; however, before I submit the article, I would like to get different perspectives on what I have written.

How can you help?

I would like you to read the attached draft of the article and let me know what you think of it. Specifically if you could answer the following questions:

- Do you think the article fairly represents general public perception?
- Is there anything missing? Or not covered in enough detail?
- Do you have any other comments?
Appendix 3.4. IJPDS publication: National data opt out programme

International Journal of Population Data Science

Journal Website: www.ipds.org

National data opt out programme: consequences for maternity statistics in England

Lewis, KM* and Hardie, P

Abstract

Electronic health records offer great potential for individual care, service improvement and, when collated, the health of the wider population. Datasets composed of these types of records have been invaluable to our understanding of risk factors for maternal and infant ill-health. However, a potential barrier to data quality in England is emerging where patients choose to opt out of sharing their information beyond the NHS. Focusing on maternity statistics, we will present the importance of population level health data for monitoring NHS services, and the potential consequences for patients of opting out. Evidencing the success of similar systems in Nordic countries, we argue that the English population must be better informed of the implications of opting out of sharing NHS data for research and the safeguards in place to protect patient information.

Background

Patient records across healthcare providers in England are being digitalised—a process whereby clinical data are stored in digital form and shared with authorised users—to the benefit of patients [1]. When compared to handwritten notes, computerised records improve the detail, completeness and reliability of patient data [1]. Electronic patient record systems improve communication between health professionals within and between different providers [2]. Patients who access their health records online report improved self-care, greater satisfaction with the communication from their doctor and, in some instances, improved safety through patient identification of medication errors [2]. The benefits of electronic health records are not limited to the care of individual patients. When collated, these data are a cost effective way to advance the health of the population through improved knowledge of healthcare services and the aetiology and treatments of health conditions [3, 4]. Electronic health records in the UK have been successfully used, for example, to investigate: neonatal impact of antibiotic prescription during pregnancy; success of different interpregnancy intervals on pregnancy following miscarriage; and pregnancy complications after cesarean section at first birth [5-7].

However, just as the use of electronic health for research and planning becomes more common place [8], new barriers to the quality of the data are also surfacing. In particular, some patients are choosing not to share their information beyond the NHS for anything other than their direct care. Opting out (see Piel et al. [9] for details about the different types of opt outs) was first made available for patients of the English NHS in January 2014 in response to a recommendation by Dame Fiona Caldicott in her 2013 information governance review [10]. This review was published amid severe concerns, voiced by both the general public and experts in the medical field, about the security of patient data following the release of individual-level health records to profit-making companies [11, 12]. A new consent model intended to be simpler and easier to access (through an online platform) launched in May 2018; however, initial figures suggest that very few patients know about this scheme [13]. As stated in the 2016 Caldicott review [14], ‘patients have a right under the NHS Constitution to request that their personal confidential information is not used beyond their direct care’. We argue that, in parallel with information about this choice, it is imperative that patients have complete and transparent information about the uses and potential advantages of sharing their information.

NHS data sharing and safeguards

The data capture organisation within the English NHS, NHS Digital, collects and stores some of the information recorded when individuals receive health or social care in England. This includes records of diagnoses and operations recorded during hospital admissions. The data are used for a variety of reasons including planning NHS services and monitoring patient safety [15]. Strictly controlled release of some patient information may be shared with NHS providers and commissioners, university researchers, charities and companies that are partnered with the NHS. Where permission is granted, all organisations must follow stringent protocols when storing and analysing the data. Personal identifiers, such as names and NHS numbers are removed in all circumstances apart from where specific
patient consent is given or where required by the law. NHS Digital states that 'we make sure data is only used for the good of health and care', and all organisations go through a lengthy application process to ensure this is the case.

**NHS maternity statistics**

The most comprehensive source of information on all births and deliveries in the NHS in England is Hospital Episode Statistics (HES), a dataset that includes all admissions to NHS or NHS-funded hospitals in England [3]. Delivery information, such as the place of delivery, baby’s sex, birthweight, gestational age and method of delivery, is used for many purposes, including to create resources for parents-to-be, to evaluate and improve maternity care provision and to investigate multiple risk factors for ill-health in mothers and babies [16]. Tools have been created utilising this data to aid expectant parents when making maternity choices. One such resource, the Birth Choices tool from Which? [17] recommends considering essential maternity statistics when planning which hospital to give birth in, such as variation in caesarean sections, induction and other medical interventions. Further, the National Maternity and Perinatal Audit (NMPA) was set up in 2016 to evaluate quality in NHS maternity services [18]. Using maternity data from HES linked with data from each maternity unit, the NMPA provides a range of statistics comparing outcomes at maternity unit level, including induction of labour and caesarean section rates. In addition, HES has been used for maternal and child health research to examine, for example, factors explaining excess child mortality in England and the safety of surgical procedures during pregnancy [19, 20]. Individual level data are necessary for this type of research, since this allows multiple risk factors for maternal and child outcomes to be taken into account.

**Opting out and the effect on the quality of NHS maternity data**

As at 1 December 2013, the average national data opt-out rate across England was 2.8% [21]. 'Top-level demographic' information published by NHS Digital shows that rates of opt-outs are higher in older people and females [21]. This is in keeping with findings from surveys and qualitative research that the characteristics of people who are less willing to share their health data with researchers differ from those who are willing to consent (although these characteristics have not been consistent across studies) [22, 23]. For example, a study of moches in the UK Millennium Cohort Study found that the proportion consenting to link survey data with their child’s NHS records differed by country of residence, age, ethnicity, lone parenthood status and education [24]. As shown in Figure 1, opt out rates in the general population are also not uniformly distributed across geographical area in England. Twelve Clinical Commissioning Groups (CCGs) have opt out rates higher than 5% and, strikingly, one CCG has a rate of 10.1% [21]. At the GP practice level, there are instances where the entire patient population have opted out [21]. As remarked by Piel et al. [9], this raises significant questions about whether the patients in these practices explicitly opted out for themselves.

To exemplify the impact of biased data on maternal information, we simulated outcome rates for two health indicators had opt outs not been applied to the data. We downloaded publically available CCG-level information from Public Health England’s Fingertips Child and Maternal Health Profiles for one common and one rare outcome, the proportion of deliveries with caesarean sections (occurring in approximately 27% of births nationwide) and births with very low birth weight (<1500g. 1.2% nationwide) [25]. We chose three CCGs for our example, each with a different rate of opt out, as published at 1st December 2013 by NHS Digital [26]. Under the assumption that the women in the maternity dataset opted out at the same rate as the whole CCG population, we modelled three scenarios based on the rate of events in the women that opted out: 1) no events; 2) events occurring at the same rate as women who did not opt out; 3) all had an event. Matching methods used by Public Health England [27]. 95% confidence intervals (CIs) were calculated in Stata 15.0 [28] using the Wilson Score method. Microsoft Excel 2013 was used to create graphs.

Table 1 displays published and simulated event rates for caesarean sections and infants born with very low birth weight for Oldham (10.1% opt outs), Mercon (2.8% opt outs) and Bradford City (0.5% opt outs) CCGs. In scenario 2, where patients opting out have the same rate of events as patients not opting out, published and simulated event rates do not differ. Figure 2 and 3 show the possible range of rates between the extremes of scenario 1 (no new events) and scenario 3 (all opt outs have an event). These scenarios show that both the CCGs with average (Mercon) and high (Oldham) rates of opting out could be showing misleading information. For the rarer event, births with very low birth weight, the rate of births with this outcome could potentially be 9-fold higher than the published rate.

The non-random nature of opt-outs has potentially large implications for the outcomes of public health monitoring, research studies and clinical audits. At an NHS trust level, information bias could produce flawed outputs in audits intending to suggest improvements and highlighting good practice. These biases will also affect the reliability of findings on maternal and children’s health research. As shown in our simulation, this is particularly the case where less common (but often more serious) outcomes are studied. Problematically, the bias introduced into datasets with opt outs applied cannot be treated with the same statistical methods used to treat missing data. Multiple imputation, a method which is commonly applied to deal with missing data, relies on the assumption that the missing information can be explained by differences in the observed data [29]. However, once patients have opted out, their data is completely removed from the dataset (i.e. complete case removal) meaning that we cannot account for systematic differences. Using multiple imputation in these instances may actually add further bias to results [29]. Methods used in population-based surveys to overcome biases of non-consent, such as weighting adjustments or simulation studies, ideally require detailed data on the population (e.g. by gender, age, deprivation level and local area) who have opted out so that correct weights can be derived - currently not published by NHS Digital. Research to determine whether those who have opted out of sharing data are different in terms of socio-demographic and health characteristics, in specific population
Figure 1: Rates of patients opting out, by CCG: England (with inlay map of London CCGs), as at December 2018. Data from NHS Digital [21].

Table 1: Published and simulated number of events, patients and event rates, by indicator and CCG. Data from the National Child and Maternal Health Intelligence Network [25] and NHS Digital [26].

<table>
<thead>
<tr>
<th>CCG opt out rate (%)</th>
<th>Bradford City CCG 0.3</th>
<th>Merthyr CCG 2.74</th>
<th>Oldham CCG 10.08</th>
<th>Bradford City CCG 0.3</th>
<th>Merthyr CCG 2.74</th>
<th>Oldham CCG 10.08</th>
</tr>
</thead>
<tbody>
<tr>
<td>Events</td>
<td>479</td>
<td>949</td>
<td>792</td>
<td>36</td>
<td>30</td>
<td>30</td>
</tr>
<tr>
<td>Deliveries</td>
<td>2024</td>
<td>3256</td>
<td>2797</td>
<td>1557</td>
<td>3219</td>
<td>3195</td>
</tr>
<tr>
<td>Event rate</td>
<td>23.6</td>
<td>21.0</td>
<td>29.6</td>
<td>21.7</td>
<td>21.2</td>
<td>21.2</td>
</tr>
<tr>
<td>(95% CI)</td>
<td>(21.8-25.5)</td>
<td>(25.3-30.6)</td>
<td>(26.7-30.0)</td>
<td>(15.7-2.99)</td>
<td>(0.89-1.65)</td>
<td>(0.89-1.65)</td>
</tr>
</tbody>
</table>

Published maternity data [25]

<table>
<thead>
<tr>
<th>New event rate (95% CI)*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Events</td>
</tr>
<tr>
<td>Deliveries</td>
</tr>
<tr>
<td>Event rate</td>
</tr>
<tr>
<td>(95% CI)</td>
</tr>
</tbody>
</table>

Scenario (1) = published events/(deliveries + extra deliveries) where extra deliveries = published deliveries/(1 - opt out rate) - published deliveries; scenario 2 (average events) = ((extra deliveries * event rate) + published events)/(deliveries + extra deliveries); scenario 3 (max. events) = (events + extra deliveries)/(deliveries + extra deliveries).

*Scenario 1 (no new events) = published events/(deliveries + extra deliveries) where extra deliveries = published deliveries/(1 - opt out rate) - published deliveries; scenario 2 (average events) = ((extra deliveries * event rate) + published events)/(deliveries + extra deliveries); scenario 3 (max. events) = (events + extra deliveries)/(deliveries + extra deliveries).
Figure 2: Observed and simulated rates of deliveries by caesarean section: By CCG, 2016/17 [21, 25]

Figure 3: Observed and simulated rates of births with very low birth weight (<1500g) observed and simulated rates: by CCG, 2016 [21, 25]
such as expectant mothers, would help in applying such methods to tackle data missing due to opt outs.

An inevitable problem?

The experience of public health researchers in the Nordic countries demonstrate that data sharing can be achieved with buy-in from citizens and at great value to clinical research [30]. In these countries, residents are assigned a personal identity number from birth, which can be used to track individuals across time and generations. Unlike the UK equivalent (NHS number), which is used only in health and social care settings [31], personal identity numbers are used across a multitude of sectors. This means that individual data can be linked across health, education and social security datasets, for example, providing high quality comprehensive information on risk factors for ill-health and other outcomes. Research using these administrative datasets have contributed markedly to the evidence base regarding determinants of health and disease across the life course. Valuable research outcomes include the long-term social and medical consequences of preterm birth and heritability of pre-eclampsia, amongst others [32, 33].

Before a data-based research project can begin, approval must be gained from a regional or national ethics committee. To protect confidentiality, personal identifiers are not shared with researchers and results cannot be published at an individual level [34]. In essence, the safeguards in place are very similar to those in England. In contrast, however, there is broad public general awareness and acceptance of the use of individual data in research and a long-standing culture of trust in public services and data donation for the good of the population [4, 30].

There is no shortage of evidence to suggest that there is no shortage of evidence to suggest that public perception of using administrative data for research between England and the Nordic countries has arisen, what can be done to improve public trust in England, and who is best placed to do it.

Evidence from England suggests that greater knowledge of research processes and safeguards improves the likelihood of acceptance of electronic health records being used without explicit consent [35]. An electronic real-time dataset integrating primary and secondary care was successfully implemented over a decade ago in Salford, Manchester. All patients were sent a letter with information and a query about opting out. Less than 0.2% of the nearly quarter of a million patients chose to opt out [36]. In contrast, information about the new withdrawn carbondioxide scheme to integrate primary and secondary care records was disseminated by generic leaflets, which were reportedly not seen by the majority of the population [11]. However, given that some of the opt-outs may have been driven by GP practices rather than patient-level decisions [9], it is not clear to what extent whether it was the NHS information campaign directed at patients (or indeed lack of it) that led to the 2.6% opt-out rate, or a lack of buy-in from clinicians.

In terms of providing information to the general public, some lessons have been learnt since care.data. A national radio campaign ran for 6-weeks after the launch of the new opt out system and NHS Digital’s website now links to Understanding Patient Data [15], an informative website run by Wellcome Trust. To further exhibit the benefits of sharing data for audits and research, NHS Digital could start by listing examples of how health care data have been used for research, as available in other NHS held datasets (e.g. CPRD [57]). Other innovative examples of dissemination include the University of Manchester’s citizen’s jury on health records [58] and an animation created as part of the #data4saveslife campaign by the Farr Institute [30]. However, the impact and wider reach of these schemes are not clear.

Arguably, information about the benefit of data sharing can only go so far in raising public confidence. Evidence from the research literature and reflected in media coverage suggests that there is unease about the potential of commercial entities, such as pharmaceutical and insurance companies, to make profit from NHS data. This is in contrast to the largely positive view of university researchers or NHS staff making use of this data [22, 36]. Therefore, Welcome Trust’s call for clear examples of “acceptable and unacceptable purposes” for which data can and cannot be used, amongst other things, should be heeded [15]. NHS Digital is beginning to advertise the benefits of sharing NHS data for research and planning purposes, and it is vital that these efforts are continued and extended. Clinicians play a pivotal role in the discourse of patient consent to use NHS data for research. Their concerns must be better understood and addressed in future consultations and information campaigns about using data for research.

Conclusion

When patients choose to opt out of sharing data beyond their direct care, the reliability of service information and evaluation and wider research based on electronic health records is diminished. We call for more transparent, clear and detailed information on: who can apply to use NHS data and for what reasons; the safeguards in place to protect individual information; and, importantly, the wider consequences of opting out on population health research and public health service information. Only with this information can individuals be expected to make an informed decision about opting out of sharing their data.

Acknowledgments

Patient involvement: We thank Maurice Hoffman, Shilpa Patel, Katherine Ruane and other representatives from the Farr Institute London Public Panel for their perspectives on the issues discussed, as well as specific comments on an earlier version of this paper.

Funding Statement

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Statement on conflicts of Interest

The authors declare that they have no conflicts of interest.
Appendix: Chapter 3


Ethics Statement

This study used open source aggregate-level data and, therefore, ethical approval was not required.

References


11. McCartney M. Care.data doesn’t care enough about consent. BMJ. 2014;348. https://doi.org/10.1136/bmj.g2631


APPENDIX - CHAPTER 4

Appendix 4.1. Thorax publication: Geospatial and seasonal variation of bronchiolitis in England
ORIGINAL RESEARCH

Geospatial and seasonal variation of bronchiolitis in England: a cohort study using hospital episode statistics

Kate Marie Lewis, Blanca De Stavola, Pia Hardelid

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Appendix 4.2. Stata code for Chapter 4 harmonic Poisson regression analyses

use "dataset\stsplit_2012to2016.dta", clear // survival set dataset

*create person-time
sts split week_pyr s, at(0(7.024038)365.25)
gen pyrs = _t-_t0

*create timeseries: from dates
gen date = _origin+_t
format date %td
gen year_event = year(date)
gen month_event = month(date)
gen week_event = week(date)
drop date

*invalid rows
drop if _st==0
drop if year_event==2011 | year_event==2017

save "geo\geo_stsplit.dta", replace

*++++++ HARMONIC REGRESSION MODEL: by REGION ++++++++ 

*convert data into easier to manage file
use "geo\geo_stsplit.dta", clear
keep week_event region year_event hesid d_entry end_fup end ///
start_bronchiolitis_any_use end_bronchiolitis_any_use ///
event_st_d_origin_t_t0 year_event
qui str ate region week_event year_event, output ///
("geo\geo_stsplit_strate region.dta", replace)

use "geo\geo_stsplit_strate region.dta", clear

*generate cos sine terms
gen sine = sin(2*_pi*1/52*(week_event-1))
gen cos = cos(2*_pi*1/52*(week_event-1))

*final model
poisson _D c.sine#ib7.region c.cos#ib7.region sine ib2013.year_event ///
ib7.region, vce(robust) exp(_Y)

*test interaction
testparm c.sine#ib7.region c.cos#ib7.region

*event prediction for duration of epidemic
predict event_region
preserve
collapse (sum) event_region _Y, by (region week_event)
gen rate=(event_region/_Y)*365250
list region week_event rate, divider clean
restore

*lincom for week 51
forvalues k=1(1)9{
    lincom `k'.region+(.9709418*cos#`k'.region)+(-.2393157*sine#`k'.region), irr
}

*SEASON PARAMETERS
gen b0=_b[_cons]
gen b1=_b[sine]
gen b2=_b[cos]
forvalues k=1(1)9{
gen alpha`k'= _b[`k'.region]
gen delta1`k'= _b[c.sine#`k'.region]
gen delta2`k'= _b[c.cos#`k'.region]
}
forvalues k=1(1)9{
gen b0`k'=b0+alpha`k'
gen b1`k'=b1+delta1`k'
gen b2`k'=b2+delta2`k'
}
forvalues k=1(1)9{
    label var b0`k' "intercept for region_`k"
    label var b1`k' "sine coef for region_`k"
    label var b2`k' "cosine coef for region_`k"
}

*amplitude, gamma: a*sqrt(b1^2+b2^2), where a=-1 if b2<0
ngen a=1
forvalues k=1(1)9{
    replace a=-1 if b2`k'<0
    gen gamma`k'=a*(sqrt(b1`k'^2+b2`k'^2))
}
forvalues k=1(1)9{
nlcom sqrt(_b[sine]+_b[sine#`k'.region])^2+(_b[cos]+_b[cos#`k'.region])^2)
}

**exp(amplitude) in poisson models
forvalues k=1(1)9{
gen gamma`k'_rate=exp(gamma`k')
}
forvalues k=1(1)9{
nlcom 
\exp(\sqrt{(_b[sine]+_b[sine#`k'.region])^2+(_b[cos]+_b[cos#`k'.region])^2})
}

*phase (in radians), psi:arctan(b1/b2)
forvalues k=1(1)9{
gen psi`k'=atan(b1`k'/b2`k')
}

forvalues k=1(1)9{
    nlcom atan((_b[sine]+_b[sine#`k'.region])/_b[cos]+_b[cos#`k'.region])
}

*estimates
forvalues k=1(1)9{
gen peak_week`k'=52.14*(psi`k'/(2*_pi))+1
}

forvalues k=1(1)9{
    nlcom 52.14* ((atan((_b[sine]+_b[sine#`k'.region]) / (_b[cos]+_b[cos#`k'.region])))/(2*_pi))+1
}

******* HARMONIC REGRESSION MODEL: by CCG **********

*convert data into easier to manage file
use "geo\geo_stsplit.dta",clear
keep week_event region CCG_number year_event hesid d_entry end_fup ///
    end_bronchiolitis_any_use end_bronchiolitis_any_use ///
    event _st _d _origin _t _t0 year_event
qui strate CCG_number week_event year_event, output ("geo\geo_stsplit_strate.dta", replace)
use "geo\geo_stsplit_strate.dta", clear

*generate cos sine terms
gen sin=sin(2*_pi*1/52.14*(week_event-1))
gen cos=cos(2*_pi*1/52.14*(week_event-1))

mepoisson _D sin cos ib2013.year_event, exp(_Y) || CCG_number: sin cos

*random effects intercept:
predict rem_sin rem_cos rem_cons, remeans
predict re_sin re_cos re_cons,reffects
gen b1=_b[sine]+re_sin
gen b2=_b[cos]+re_cos
gen b0=_b[_cons]+re_cons
gen IR_CCG=rem_cons

*SEASON PARAMETERS

*amplitude, gamma: \( a \cdot \sqrt{b_1^2+b_2^2} \), where \( a = -1 \) if \( b_2 < 0 \)

```stata
gen a = 1
replace a = -1 if b2 < 0
gen gamma = a * (sqrt(b1^2 + b2^2))
```

**exp(amplitude) in poisson models

```stata
gen gamma_rate = exp(gamma)
```

*phase (in radians), psi: \( \arctan(b_1/b_2) \)

```stata
gen psi = atan(b1/b2)
```

*estimates

```stata
gen peak_week = 52.14 * (psi / (2 * _pi)) + 1
gen mean_rate = exp(b0) * 365250
```
### Appendix 4.3. Harmonic Poisson regression models

**Table A4.1.** Output from the harmonic Poisson regression model for rates of bronchiolitis, by region and year

<table>
<thead>
<tr>
<th>Region</th>
<th>Coefficient</th>
<th>p</th>
<th>(95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>North East</td>
<td>0.51</td>
<td>&lt;0.001</td>
<td>(0.47, 0.56)</td>
</tr>
<tr>
<td>North West</td>
<td>0.68</td>
<td>&lt;0.001</td>
<td>(0.65, 0.71)</td>
</tr>
<tr>
<td>Yorkshire &amp; the Humber</td>
<td>0.49</td>
<td>&lt;0.001</td>
<td>(0.45, 0.52)</td>
</tr>
<tr>
<td>East Midlands</td>
<td>0.31</td>
<td>&lt;0.001</td>
<td>(0.27, 0.35)</td>
</tr>
<tr>
<td>West Midlands</td>
<td>0.50</td>
<td>&lt;0.001</td>
<td>(0.47, 0.54)</td>
</tr>
<tr>
<td>East of England</td>
<td>0.16</td>
<td>&lt;0.001</td>
<td>(0.12, 0.19)</td>
</tr>
<tr>
<td>London Reference</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>South East</td>
<td>0.26</td>
<td>&lt;0.001</td>
<td>(0.23, 0.29)</td>
</tr>
<tr>
<td>South West</td>
<td>0.36</td>
<td>&lt;0.001</td>
<td>(0.32, 0.39)</td>
</tr>
<tr>
<td>Sine</td>
<td>-0.41</td>
<td>&lt;0.001</td>
<td>(-0.43, -0.39)</td>
</tr>
<tr>
<td>Cosine</td>
<td>1.16</td>
<td>&lt;0.001</td>
<td>(1.13, 1.18)</td>
</tr>
<tr>
<td>Sine* Region North East</td>
<td>0.07</td>
<td>&lt;0.001</td>
<td>(0.03, 0.11)</td>
</tr>
<tr>
<td>North West</td>
<td>-0.07</td>
<td>&lt;0.001</td>
<td>(-0.10, -0.04)</td>
</tr>
<tr>
<td>Yorkshire &amp; the Humber</td>
<td>-0.01</td>
<td>0.46</td>
<td>(-0.05, 0.02)</td>
</tr>
<tr>
<td>East Midlands</td>
<td>0.03</td>
<td>0.15</td>
<td>(-0.01, 0.07)</td>
</tr>
<tr>
<td>West Midlands</td>
<td>-0.04</td>
<td>0.03</td>
<td>(-0.07, 0.00)</td>
</tr>
<tr>
<td>East of England</td>
<td>-0.03</td>
<td>0.16</td>
<td>(-0.06, 0.01)</td>
</tr>
<tr>
<td>London Reference</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>South East</td>
<td>-0.06</td>
<td>&lt;0.001</td>
<td>(-0.09, -0.02)</td>
</tr>
<tr>
<td>South West</td>
<td>0.06</td>
<td>&lt;0.001</td>
<td>(0.02, 0.09)</td>
</tr>
<tr>
<td>Cosine* Region North East</td>
<td>0.41</td>
<td>&lt;0.001</td>
<td>(0.36, 0.46)</td>
</tr>
<tr>
<td>North West</td>
<td>0.21</td>
<td>&lt;0.001</td>
<td>(0.18, 0.24)</td>
</tr>
<tr>
<td>Yorkshire &amp; the Humber</td>
<td>0.25</td>
<td>&lt;0.001</td>
<td>(0.21, 0.29)</td>
</tr>
<tr>
<td>East Midlands</td>
<td>0.32</td>
<td>&lt;0.001</td>
<td>(0.28, 0.37)</td>
</tr>
<tr>
<td>West Midlands</td>
<td>0.23</td>
<td>&lt;0.001</td>
<td>(0.19, 0.27)</td>
</tr>
<tr>
<td>East of England</td>
<td>0.37</td>
<td>&lt;0.001</td>
<td>(0.32, 0.41)</td>
</tr>
<tr>
<td>London Reference</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>South East</td>
<td>0.28</td>
<td>&lt;0.001</td>
<td>(0.25, 0.32)</td>
</tr>
<tr>
<td>South West</td>
<td>0.32</td>
<td>&lt;0.001</td>
<td>(0.28, 0.37)</td>
</tr>
<tr>
<td>Year of event</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2012</td>
<td>0.06</td>
<td>&lt;0.001</td>
<td>(0.05, 0.08)</td>
</tr>
<tr>
<td>2013</td>
<td></td>
<td></td>
<td>Reference</td>
</tr>
<tr>
<td>2014</td>
<td>0.06</td>
<td>&lt;0.001</td>
<td>(0.04, 0.07)</td>
</tr>
<tr>
<td>2015</td>
<td>0.22</td>
<td>&lt;0.001</td>
<td>(0.20, 0.23)</td>
</tr>
<tr>
<td>2016</td>
<td>0.27</td>
<td>&lt;0.001</td>
<td>(0.25, 0.30)</td>
</tr>
<tr>
<td>Constant</td>
<td>-9.85</td>
<td>&lt;0.001</td>
<td>(-9.88, -9.83)</td>
</tr>
</tbody>
</table>

Wald test testing interaction between region and sine and cosine functions, p<0.0001; likelihood ratio test, p<0.0001
Table A8.2. Output of the harmonic multilevel mixed-effect Poisson regression analysis for rates of bronchiolitis

<table>
<thead>
<tr>
<th></th>
<th>Coefficient</th>
<th>(95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Individual-level</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sine</td>
<td>-0.44</td>
<td>(-0.46, -0.42)</td>
</tr>
<tr>
<td>Cos</td>
<td>1.42</td>
<td>(1.39, 1.45)</td>
</tr>
<tr>
<td>Year of event</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2012</td>
<td>0.06</td>
<td>(0.04, 0.07)</td>
</tr>
<tr>
<td>2013</td>
<td>Ref.</td>
<td></td>
</tr>
<tr>
<td>2014</td>
<td>0.06</td>
<td>(0.04, 0.08)</td>
</tr>
<tr>
<td>2015</td>
<td>0.22</td>
<td>(0.20, 0.23)</td>
</tr>
<tr>
<td>2016</td>
<td>0.27</td>
<td>(0.25, 0.28)</td>
</tr>
<tr>
<td>Constant</td>
<td>-9.54</td>
<td>(-9.58, -9.49)</td>
</tr>
<tr>
<td><strong>CCG-level</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>var(sine)</td>
<td>0.01</td>
<td>(0.01, 0.02)</td>
</tr>
<tr>
<td>var(cos)</td>
<td>0.05</td>
<td>(0.04, 0.06)</td>
</tr>
<tr>
<td>var(constant)</td>
<td>0.11</td>
<td>(0.10, 0.13)</td>
</tr>
</tbody>
</table>

Excluding London-based CCGs

Table A4.3. Harmonic multilevel mixed-effect Poisson regression analysis for rates of RSV bronchiolitis: Excluding London-based CCGs

<table>
<thead>
<tr>
<th></th>
<th>Coefficient</th>
<th>p</th>
<th>(95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Individual-level</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sine</td>
<td>-0.44</td>
<td>&lt;0.001</td>
<td>(-0.46, -0.42)</td>
</tr>
<tr>
<td>Cos</td>
<td>1.48</td>
<td>&lt;0.001</td>
<td>(1.45, 1.50)</td>
</tr>
<tr>
<td>Year of event</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2012</td>
<td>0.07</td>
<td>&lt;0.001</td>
<td>(0.05, 0.09)</td>
</tr>
<tr>
<td>2013</td>
<td>Ref.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2014</td>
<td>0.06</td>
<td>&lt;0.001</td>
<td>(0.05, 0.08)</td>
</tr>
<tr>
<td>2015</td>
<td>0.22</td>
<td>&lt;0.001</td>
<td>(0.20, 0.24)</td>
</tr>
<tr>
<td>2016</td>
<td>0.27</td>
<td>&lt;0.001</td>
<td>(0.25, 0.28)</td>
</tr>
<tr>
<td>Constant</td>
<td>-9.44</td>
<td>&lt;0.001</td>
<td>(-9.49, -9.40)</td>
</tr>
<tr>
<td><strong>CCG-level</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>var(sine)</td>
<td>0.01</td>
<td></td>
<td>(0.01, 0.02)</td>
</tr>
<tr>
<td>var(cos)</td>
<td>0.03</td>
<td></td>
<td>(0.03, 0.04)</td>
</tr>
<tr>
<td>var(constant)</td>
<td>0.10</td>
<td></td>
<td>(0.08, 0.13)</td>
</tr>
</tbody>
</table>
Appendix 4.4: Sensitivity analyses

Bootstrapped standard errors (SEs)

Table A4.4. Multivariable linear regression analyses assessing the effect of population density and IMD score on IRR and peak timing at the CCG-level following two stage analysis with 1,000 bootstraps on a 20% sample of model

<table>
<thead>
<tr>
<th></th>
<th>No bootstrapped SEs</th>
<th>1000 bootstrapped SEs</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Coefficient Lower CI Upper CI</td>
<td>Coefficient Lower CI Upper CI</td>
</tr>
<tr>
<td><strong>IRR</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Population density*</td>
<td>-0.1046 -0.1375 -0.0718</td>
<td>-0.1046 -0.1410 -0.0683</td>
</tr>
<tr>
<td>IMD</td>
<td>0.0214 0.0160 0.0268</td>
<td>0.0214 0.0159 0.0269</td>
</tr>
<tr>
<td><strong>Peak week</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Population density*</td>
<td>-0.1471 -0.1532 -0.1411</td>
<td>-0.1471 -0.2288 -0.0655</td>
</tr>
<tr>
<td>IMD</td>
<td>0.0496 0.0457 0.0534</td>
<td>0.0496 -0.0079 0.1070</td>
</tr>
<tr>
<td>IMD^2</td>
<td>-0.0007 -0.0007 -0.0006</td>
<td>-0.0007 -0.0018 0.0004</td>
</tr>
</tbody>
</table>

*Log-scale

Note: Year of admission was excluded from model due to convergence problem

One admission per child

Table A4.5. Derived average annual seasonal estimates following harmonic Poisson regression (not shown)*, by region: England, 2012 to 2016 (one admission per child)

<table>
<thead>
<tr>
<th>Region*</th>
<th>N</th>
<th>Admissions Rate per 1000 infant-years (95% CI)</th>
<th>Amplitude $\hat{\gamma}^{**}$ (95% CI)</th>
<th>Peak week $\bar{p}$ (95%CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>North East</td>
<td>8,471</td>
<td>57.8 (56.6, 59.1)</td>
<td>5.66 (4.91, 6.52)</td>
<td>51.3 (50.8, 51.7)</td>
</tr>
<tr>
<td>North West</td>
<td>20,017</td>
<td>61.2 (60.4, 61.9)</td>
<td>4.86 (4.30, 5.49)</td>
<td>50.1 (49.7, 50.6)</td>
</tr>
<tr>
<td>Yorkshire &amp; the Humber</td>
<td>15,163</td>
<td>51.6 (50.7, 52.4)</td>
<td>4.88 (4.28, 5.56)</td>
<td>50.5 (50.0, 51.0)</td>
</tr>
<tr>
<td>East Midlands</td>
<td>10,939</td>
<td>45.4 (44.6, 46.3)</td>
<td>5.23 (4.57, 5.98)</td>
<td>50.9 (50.4, 51.4)</td>
</tr>
<tr>
<td>West Midlands</td>
<td>17,155</td>
<td>51.4 (50.6, 52.2)</td>
<td>4.97 (4.36, 5.67)</td>
<td>50.3 (49.9, 50.8)</td>
</tr>
<tr>
<td>East of England</td>
<td>13,543</td>
<td>40.3 (39.6, 41.0)</td>
<td>5.56 (4.83, 6.40)</td>
<td>50.6 (50.2, 51.1)</td>
</tr>
<tr>
<td>London</td>
<td>16,144</td>
<td>27.5 (27.0, 27.9)</td>
<td>3.92 (3.47, 4.42)</td>
<td>50.0 (49.5, 50.6)</td>
</tr>
<tr>
<td>South East</td>
<td>20,232</td>
<td>42.2 (41.6, 42.8)</td>
<td>5.24 (4.55, 6.04)</td>
<td>50.3 (49.8, 50.8)</td>
</tr>
<tr>
<td>South West</td>
<td>12,868</td>
<td>46.8 (46.0, 47.6)</td>
<td>5.43 (4.73, 6.23)</td>
<td>50.9 (50.5, 51.4)</td>
</tr>
<tr>
<td>**Phase shift $\bar{\psi}$ (95%CI)</td>
<td>**Duration (weeks)</td>
<td>**IRR at week 51 (95% CI) **</td>
<td></td>
<td></td>
</tr>
<tr>
<td>North East</td>
<td>-0.23 (-0.29, -0.17)</td>
<td>26 (26-52)</td>
<td>2.46 (2.11, 2.87)</td>
<td></td>
</tr>
<tr>
<td>North West</td>
<td>-0.36 (-0.42, -0.31)</td>
<td>26 (25-51)</td>
<td>2.43 (2.10, 2.82)</td>
<td></td>
</tr>
<tr>
<td>Yorkshire &amp; the Humber</td>
<td>-0.32 (-0.38, -0.26)</td>
<td>26 (25-51)</td>
<td>2.06 (1.78, 2.40)</td>
<td></td>
</tr>
<tr>
<td>East Midlands</td>
<td>-0.27 (-0.33, -0.21)</td>
<td>26 (26-52)</td>
<td>1.88 (1.61, 2.18)</td>
<td></td>
</tr>
<tr>
<td>West Midlands</td>
<td>-0.34 (-0.40, -0.29)</td>
<td>26 (25-51)</td>
<td>2.07 (1.78, 2.40)</td>
<td></td>
</tr>
<tr>
<td>East of England</td>
<td>-0.30 (-0.36, -0.25)</td>
<td>26 (26-52)</td>
<td>1.69 (1.46, 1.97)</td>
<td></td>
</tr>
<tr>
<td>London</td>
<td>-0.37 (-0.43, -0.31)</td>
<td>26 (25-51)</td>
<td>Reference</td>
<td></td>
</tr>
<tr>
<td>South East</td>
<td>-0.34 (-0.40, -0.28)</td>
<td>26 (25-51)</td>
<td>1.73 (1.49, 2.02)</td>
<td></td>
</tr>
<tr>
<td>South West</td>
<td>-0.26 (-0.32, -0.21)</td>
<td>26 (26-52)</td>
<td>1.96 (1.68, 2.29)</td>
<td></td>
</tr>
</tbody>
</table>
*Adjusted for year of admission, **amplitude exponentiated

**Table A4.6.** Harmonic multilevel mixed-effect Poisson regression analysis for rates of RSV bronchiolitis (one admission per child)

<table>
<thead>
<tr>
<th></th>
<th>Coefficient</th>
<th>p</th>
<th>(95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Individual-level</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sine</td>
<td>-0.53</td>
<td>&lt;0.001</td>
<td>(-0.55, -0.51)</td>
</tr>
<tr>
<td>Cos</td>
<td>1.54</td>
<td>&lt;0.001</td>
<td>(1.51, 1.57)</td>
</tr>
<tr>
<td>Constant</td>
<td>-9.60</td>
<td>&lt;0.001</td>
<td>(-9.64, -9.56)</td>
</tr>
<tr>
<td>var(sine)</td>
<td>0.02</td>
<td></td>
<td>(0.01, 0.02)</td>
</tr>
<tr>
<td>var(cos)</td>
<td>0.05</td>
<td></td>
<td>(0.04, 0.06)</td>
</tr>
<tr>
<td>var(constant)</td>
<td>0.10</td>
<td></td>
<td>(0.09, 0.12)</td>
</tr>
</tbody>
</table>

**Note:** Year of admission was excluded from model due to convergence problems

**Table A4.7.** Multivariable linear regression analyses assessing the effect of population density and IMD on IRR compared the national average and peak timing of CCG seasonal epidemic (one admission per child)

<table>
<thead>
<tr>
<th></th>
<th>All CCGs</th>
<th></th>
<th></th>
<th>Partial $\eta^2$ (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Coefficient (95% CI)</td>
<td></td>
<td></td>
<td>Total $\eta^2$</td>
</tr>
<tr>
<td><strong>IRR</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Population density*</td>
<td>-0.09 (-0.12, -0.06)</td>
<td>0.13 (0.06, 0.22)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>IMD</td>
<td>0.02 (0.02, 0.03)</td>
<td>0.23 (0.14, 0.32)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Constant</td>
<td>1.22 (1.02, 1.42)</td>
<td></td>
<td></td>
<td>Total $\eta^2$ 0.24 (0.14, 0.34)</td>
</tr>
<tr>
<td><strong>Peak week</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Population density*</td>
<td>-0.26 (-0.31, -0.21)</td>
<td>0.32 (0.22, 0.41)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>IMD</td>
<td>0.11 (0.08, 0.15)</td>
<td>0.19 (0.10, 0.28)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>IMD$^2$</td>
<td></td>
<td></td>
<td></td>
<td>0.12 (0.05, 0.21)</td>
</tr>
<tr>
<td>Constant</td>
<td>50.59 (50.11, 51.06)</td>
<td></td>
<td></td>
<td>Total $\eta^2$ 0.40 (0.29, 0.48)</td>
</tr>
</tbody>
</table>

*Log-scale
APPENDIX - CHAPTER 5

Appendix 5.1. JECH publication: Is socioeconomic position associated with bronchiolitis seasonality?
Is socioeconomic position associated with bronchiolitis seasonality? A cohort study

Kate Lewis, Bianca De Stavola, Pia Hardelid

Supplemental material is published online only. To view please visit the journal online (http://nicobil.org/10.1136/ jeb-2019-218651).

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Appendix 5.2. Stata code for Chapter 5 harmonic regression analyses

use "dataset\stsplit_2012to2016_2.dta", clear // survival set dataset

*create person-time
stsplits week_pyrs, at(0(7.024038)365.25)
stsplits ageband, at(0, 91.3125, 182.625, 365.25)

*create age var
gen age=2 if ageband!.
replace age=1 if ageband<180
replace age=0 if ageband==0
label variable ageband "<3/4-<6/6-<12 months"

*create timeseries: from dates
gen date=_origin+_t
format date %td
gen year_event=year(date)
gen month_event=month(date)
gen week_event=week(date)
drop date

*invalid rows
drop if _st==0
drop if year_event==2011 | year_event==2017
gen pyrs=_t-_t0

*++++++ HARMONIC REGRESSION MODEL +++++++

*generate harmonics
gen sine=sin(2*_pi*1/52.14*(week_event-1))
gen cosine=cos(2*_pi*1/52.14*(week_event-1))

*final model
poisson _d ib1.mdob ib1.sex ib2013.year_event ib7.region ib2.age c.sine c.cosine ///
c.sine#i.IMD c.cosine#i.IMD c.sine#i.age c.cosine#i.age i.IMD, ///
exp(pyrs) vce(cluster hesid) irr
testparm c.sine#i.IMD c.cosine#i.IMD
testparm c.sine#i.age c.cosine#i.age

predict event_IMD
estat ic

*******************************************************

*duration of epidemic by IMD
preserve
collapse (sum) event_IMD_d pyrs, by (IMD week_event)
gen rate_obs=(d/pyrs)*365250
gen rate_pre=(event_IMD/pyrs)*365250
list IMD week_event rate_obs rate_pre, divider clean
restore

*lincom for week 50
forvalues k=1(1)5{
  lincom `k'.IMD+(.929261*\cosine#`k'.IMD)+(-.3694239*\sine#`k'.IMD),irr
}

*lincom for week 51
forvalues k=1(1)5{
  lincom `k'.IMD+(.9669321*\cosine#`k'.IMD)+(-.2550341*\sine#`k'.IMD),irr
}

*SEASON PARAMETERS
gen b01=_b[_cons]
gen b11=_b[\sine]
gen b21=_b[\cosine]
forvalues k=2(1)5{
gen alpha`k'=_b[\`k'.IMD]
gen delta1`k'=_b[\`k'.IMD#c.sine]
gen delta2`k'=_b[\`k'.IMD#c.cosine]
}
forvalues k=2(1)5{
gen b0`k'=b01+alpha`k'
gen b1`k'=b11+delta1`k'
gen b2`k'=b21+delta2`k'
}
forvalues k=1(1)5{
  label var b0`k' "intercept for IMD_`k"
  label var b1`k' "sine coef for IMD_`k"
  label var b2`k' "cosine coef for IMD_`k"
}

*amplitude, gamma: a*sqrt(b1^2+b2^2), where a=-1 if b2<0
gen a=1
forvalues k=1(1)5{
  replace a=-1 if b2`k'<0
  gen gamma`k'=a*(sqrt(b1`k'^2+b2`k'^2))
}
forvalues k=1(1)5{
\begin{verbatim}

nlcom \sqrt{(_b[sine]+_b[sine#k'.IMD])^2+(_b[cosine]+_b[cosine#k'.IMD])^2}

**exp(amplitude) in poisson models
forvalues k=1(1)5{
gen gamma_rate`k'=exp(gamma`k')
}
forvalues k=1(1)5{
  nlcom \exp(\sqrt{(_b[sine]+_b[sine#k'.IMD])^2+(_b[cosine]+_b[cosine#k'.IMD])^2})
}

*phase (in radians), psi: \texttt{-arctan(b1/b2)}
forvalues k=1(1)5{
gen psi`k'=\texttt{-atan(b1`k'/b2`k')}
}
forvalues k=1(1)5{
  nlcom \texttt{atan((_b[sine]+_b[sine#k'.IMD])/(_b[cosine]+_b[cosine#k'.IMD]))}
}

*estimates
forvalues k=1(1)5{
gen peak_week`k'=52.14*(psi`k'/(2*_pi))+1
}
forvalues k=1(1)5{
  nlcom 52.14*((atan((_b[sine]+_b[sine#k'.IMD])/(_b[cosine]+_b[cosine#k'.IMD])))/(2*_pi)))+1

\end{verbatim}
### Appendix 5.3. Harmonic Poisson regression models

#### Table A5.1. IRRs (95% CI) from multivariable harmonic Poisson regression models using IMD quintiles including and excluding the health deprivation and disability domain

<table>
<thead>
<tr>
<th>Variable</th>
<th>Full IMD† IRR (95% CI)</th>
<th>IMD excluding health domain‡ IRR (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>IMD quintile</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 - Least deprived</td>
<td>Reference</td>
<td>Reference</td>
</tr>
<tr>
<td>2</td>
<td>1.10 (1.06, 1.14)</td>
<td>1.10 (1.06, 1.14)</td>
</tr>
<tr>
<td>3</td>
<td>1.21 (1.17, 1.25)</td>
<td>1.21 (1.17, 1.26)</td>
</tr>
<tr>
<td>4</td>
<td>1.37 (1.32, 1.42)</td>
<td>1.34 (1.30, 1.39)</td>
</tr>
<tr>
<td>5 - Most deprived</td>
<td>1.62 (1.57, 1.67)</td>
<td>1.62 (1.57, 1.67)</td>
</tr>
<tr>
<td>Sine</td>
<td>0.55 (0.53, 0.56)</td>
<td>0.55 (0.53, 0.56)</td>
</tr>
<tr>
<td>Cosine</td>
<td>3.45 (3.33, 3.57)</td>
<td>3.45 (3.33, 3.57)</td>
</tr>
<tr>
<td>Age group</td>
<td></td>
<td></td>
</tr>
<tr>
<td>&lt; 3 months</td>
<td>1.85 (1.81, 1.89)</td>
<td>1.85 (0.00, 0.00)</td>
</tr>
<tr>
<td>3 to &lt; 6 months</td>
<td>1.68 (1.65, 1.72)</td>
<td>1.68 (1.81, 1.89)</td>
</tr>
<tr>
<td>6 to &lt; 12 months</td>
<td>Reference</td>
<td>Reference</td>
</tr>
<tr>
<td>Sex</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Male</td>
<td>1.46 (1.44, 1.47)</td>
<td>1.46 (1.44, 1.47)</td>
</tr>
<tr>
<td>Year of event</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2012</td>
<td>1.06 (1.04, 1.08)</td>
<td>1.06 (1.04, 1.08)</td>
</tr>
<tr>
<td>2013</td>
<td>Reference</td>
<td>Reference</td>
</tr>
<tr>
<td>2014</td>
<td>1.06 (1.04, 1.08)</td>
<td>1.06 (1.04, 1.08)</td>
</tr>
<tr>
<td>2015</td>
<td>1.24 (1.22, 1.26)</td>
<td>1.24 (1.22, 1.26)</td>
</tr>
<tr>
<td>2016</td>
<td>1.31 (1.29, 1.33)</td>
<td>1.31 (1.29, 1.33)</td>
</tr>
<tr>
<td>Region</td>
<td></td>
<td></td>
</tr>
<tr>
<td>North East</td>
<td>2.04 (1.98, 2.09)</td>
<td>2.08 (2.03, 2.14)</td>
</tr>
<tr>
<td>North West</td>
<td>2.17 (2.13, 2.22)</td>
<td>2.22 (2.17, 2.27)</td>
</tr>
<tr>
<td>Yorkshire and the Humber</td>
<td>1.82 (1.78, 1.87)</td>
<td>1.85 (1.80, 1.89)</td>
</tr>
<tr>
<td>East Midlands</td>
<td>1.66 (1.62, 1.70)</td>
<td>1.68 (1.64, 1.72)</td>
</tr>
<tr>
<td>West Midlands</td>
<td>1.84 (1.80, 1.88)</td>
<td>1.86 (1.82, 1.90)</td>
</tr>
<tr>
<td>East of England</td>
<td>1.53 (1.49, 1.57)</td>
<td>1.54 (1.50, 1.57)</td>
</tr>
<tr>
<td>London</td>
<td>Reference</td>
<td>Reference</td>
</tr>
<tr>
<td>South East</td>
<td>1.66 (1.63, 1.70)</td>
<td>1.67 (1.63, 1.71)</td>
</tr>
<tr>
<td>South West</td>
<td>1.79 (1.74, 1.83)</td>
<td>1.80 (1.76, 1.84)</td>
</tr>
<tr>
<td>Month of birth</td>
<td></td>
<td></td>
</tr>
<tr>
<td>January</td>
<td>Reference</td>
<td>Reference</td>
</tr>
<tr>
<td>February</td>
<td>1.12 (1.08, 1.16)</td>
<td>1.12 (1.08, 1.16)</td>
</tr>
<tr>
<td>March</td>
<td>1.22 (1.18, 1.26)</td>
<td>1.22 (1.18, 1.26)</td>
</tr>
<tr>
<td>April</td>
<td>1.30 (1.26, 1.35)</td>
<td>1.30 (1.26, 1.35)</td>
</tr>
<tr>
<td>May</td>
<td>1.35 (1.31, 1.40)</td>
<td>1.35 (1.31, 1.40)</td>
</tr>
<tr>
<td>June</td>
<td>1.37 (1.33, 1.42)</td>
<td>1.37 (1.33, 1.42)</td>
</tr>
<tr>
<td>July</td>
<td>1.42 (1.37, 1.47)</td>
<td>1.42 (1.37, 1.47)</td>
</tr>
<tr>
<td>August</td>
<td>1.48 (1.43, 1.53)</td>
<td>1.48 (1.43, 1.53)</td>
</tr>
<tr>
<td>September</td>
<td>1.52 (1.47, 1.57)</td>
<td>1.52 (1.47, 1.57)</td>
</tr>
<tr>
<td>October</td>
<td>1.54 (1.49, 1.58)</td>
<td>1.54 (1.49, 1.58)</td>
</tr>
<tr>
<td>November</td>
<td>1.34 (1.30, 1.38)</td>
<td>1.34 (1.30, 1.38)</td>
</tr>
<tr>
<td>December</td>
<td>1.03 (1.00, 1.06)</td>
<td>1.03 (1.00, 1.06)</td>
</tr>
<tr>
<td>Sine*age</td>
<td></td>
<td></td>
</tr>
<tr>
<td>&lt; 3 months</td>
<td>1.15 (1.12, 1.18)</td>
<td>1.15 (1.12, 1.18)</td>
</tr>
<tr>
<td>3 to &lt; 6 months</td>
<td>1.12 (1.09, 1.16)</td>
<td>1.12 (1.09, 1.16)</td>
</tr>
<tr>
<td>6 to &lt; 12 months</td>
<td>Reference</td>
<td>Reference</td>
</tr>
<tr>
<td>Cosine*age</td>
<td></td>
<td></td>
</tr>
<tr>
<td>&lt; 3 months</td>
<td>1.65 (1.61, 1.70)</td>
<td>1.65 (1.61, 1.70)</td>
</tr>
<tr>
<td>3 to &lt; 6 months</td>
<td>1.04 (1.00, 1.07)</td>
<td>1.04 (1.00, 1.07)</td>
</tr>
<tr>
<td>6 to &lt; 12 months</td>
<td>Reference</td>
<td>Reference</td>
</tr>
<tr>
<td>Sine*IMD quintile</td>
<td>1 - Least deprived</td>
<td>Reference</td>
</tr>
<tr>
<td>2</td>
<td>1.07 (1.03, 1.11)</td>
<td>1.06 (1.02, 1.10)</td>
</tr>
<tr>
<td>3</td>
<td>1.14 (1.10, 1.18)</td>
<td>1.14 (1.11, 1.18)</td>
</tr>
<tr>
<td>4</td>
<td>1.18 (1.14, 1.22)</td>
<td>1.17 (1.13, 1.21)</td>
</tr>
<tr>
<td>5 - Most deprived</td>
<td>1.21 (1.18, 1.25)</td>
<td>1.21 (1.18, 1.25)</td>
</tr>
<tr>
<td>Cosine*IMD quintile</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 - Least deprived</td>
<td>Reference</td>
<td>Reference</td>
</tr>
<tr>
<td></td>
<td>Wald test of interaction between age and sine and cosine functions, $\chi^2(8)=239.2$, $\rho&lt;0.0001$</td>
<td>Wald test of interaction between IMD quintiles and sine and cosine functions, $\chi^2(4)=1835.9$, $\rho&lt;0.0001$; †Wald test of interaction between age and sine and cosine functions, $\chi^2(8)=253.8$, $\rho&lt;0.0001$, Wald test of interaction between IMD quintiles and sine and cosine functions, $\chi^2(4)=1836.0$, $\rho&lt;0.0001$</td>
</tr>
<tr>
<td>---</td>
<td>---</td>
<td>---</td>
</tr>
<tr>
<td>2</td>
<td>0.99 (0.95, 1.04)</td>
<td>1.00 (0.95, 1.04)</td>
</tr>
<tr>
<td>3</td>
<td>0.99 (0.95, 1.04)</td>
<td>0.99 (0.95, 1.03)</td>
</tr>
<tr>
<td>4</td>
<td>0.95 (0.91, 0.98)</td>
<td>0.96 (0.93, 1.00)</td>
</tr>
<tr>
<td>5 - Most deprived</td>
<td>0.90 (0.87, 0.94)</td>
<td>0.90 (0.87, 0.93)</td>
</tr>
</tbody>
</table>
## APPENDIX - CHAPTER 6

### Appendix 6.1. ESC-DAG output

Table A6.1. Study decision logs

<table>
<thead>
<tr>
<th><strong>Publication details #1</strong>&lt;sup&gt;103&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Author</strong></td>
</tr>
<tr>
<td><strong>Year</strong></td>
</tr>
<tr>
<td><strong>Study title</strong></td>
</tr>
<tr>
<td><strong>Design</strong></td>
</tr>
<tr>
<td><strong>Methods</strong></td>
</tr>
<tr>
<td><strong>Sample</strong></td>
</tr>
<tr>
<td><strong>Variables</strong></td>
</tr>
<tr>
<td><strong>Outcome</strong></td>
</tr>
<tr>
<td><strong>Study exposure</strong></td>
</tr>
<tr>
<td><strong>My exposure of interest</strong></td>
</tr>
<tr>
<td><strong>Control(s)</strong></td>
</tr>
<tr>
<td><strong>Mediator(s)</strong></td>
</tr>
<tr>
<td><strong>Moderator(s)</strong></td>
</tr>
<tr>
<td><strong>Excluded</strong></td>
</tr>
</tbody>
</table>

### Diagrams

**Implied graph**<sup>*</sup>

![Implied graph](image1)

**DAG**<sup>*</sup>

![DAG](image2)
### Publication details #2

<table>
<thead>
<tr>
<th>Author</th>
<th>Hannah C Moore, Nicholas de Klerk, Christopher C Blyth, et al.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Year</td>
<td>2019</td>
</tr>
<tr>
<td><strong>Study title</strong></td>
<td>Temporal trends and socioeconomic differences in acute respiratory infection hospitalisations in children: an intercountry comparison of birth cohort studies in Western Australia, England and Scotland</td>
</tr>
<tr>
<td><strong>Design</strong></td>
<td>Retrospective population-based cohort studies: Linked birth, death and hospitalisation data</td>
</tr>
<tr>
<td><strong>Methods</strong></td>
<td>Negative binomial regression models</td>
</tr>
<tr>
<td><strong>Sample</strong></td>
<td>6,976,508 children 0-4 years</td>
</tr>
<tr>
<td><strong>Variables</strong></td>
<td></td>
</tr>
<tr>
<td><strong>Outcome</strong></td>
<td>Bronchiolitis hospitalisations</td>
</tr>
<tr>
<td><strong>Exposure(s)</strong></td>
<td>Socioeconomic deprivation; year</td>
</tr>
<tr>
<td><strong>Control(s)</strong></td>
<td>sex</td>
</tr>
<tr>
<td><strong>Mediator(s)</strong></td>
<td>-</td>
</tr>
<tr>
<td><strong>Moderator(s)</strong></td>
<td>-</td>
</tr>
<tr>
<td><strong>Excluded</strong></td>
<td>-</td>
</tr>
</tbody>
</table>

**ALL EDGES HAVE ALREADY BEEN ASSESSED**

### Publication details #3

<table>
<thead>
<tr>
<th>Author</th>
<th>Lawder et al.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Year</td>
<td>2019</td>
</tr>
<tr>
<td><strong>Study title</strong></td>
<td>Impact of maternal smoking on early childhood health: a retrospective cohort linked dataset analysis of 697 003 children born in Scotland 1997–2009</td>
</tr>
<tr>
<td><strong>Design</strong></td>
<td>Retrospective population-based cohort study; Linked birth, death, maternity, infant health, child health surveillance and admission records</td>
</tr>
<tr>
<td><strong>Methods</strong></td>
<td>Cox and logistic regression. Population attributable fraction (PAF)</td>
</tr>
<tr>
<td><strong>Sample</strong></td>
<td>Singleton births between 1997 and 2009 (n=697 003) followed to March 2012.</td>
</tr>
<tr>
<td><strong>Variables</strong></td>
<td></td>
</tr>
<tr>
<td><strong>Outcome</strong></td>
<td>Hospital admissions with a primary diagnosis of bronchiolitis</td>
</tr>
<tr>
<td><strong>Exposure(s)</strong></td>
<td>Smoking during pregnancy</td>
</tr>
<tr>
<td><strong>Control(s)</strong></td>
<td>Scottish Index of Multiple Deprivation (SIMD), maternal age, infant gender, parity, delivery method, infant breastfed at 6–8 weeks, mother’s country of birth</td>
</tr>
<tr>
<td><strong>Mediator(s)</strong></td>
<td>-</td>
</tr>
<tr>
<td><strong>Moderator(s)</strong></td>
<td>-</td>
</tr>
<tr>
<td><strong>Excluded/changed</strong></td>
<td>Mother’s socioeconomic status and father’s socioeconomic status. Reason: high correlation with SIMD.</td>
</tr>
<tr>
<td><strong>Diagrams</strong></td>
<td></td>
</tr>
</tbody>
</table>
Publication details #4

**Author**  Kennedy et al.
**Year**  2018
**Study title**  Associations of mobile source air pollution during the first year of life with childhood pneumonia, bronchiolitis, and otitis media
**Design**  Birth cohort; Kaiser Air Pollution and Pediatric Asthma Study, a retrospective birth cohort of children born during 2000–2010 and insured by Kaiser Permanente Georgia (USA)
**Methods**  Cox proportional hazards models
**Sample**  22,441 children followed until 2 years
**Variables**

**Outcome**  Diagnosis of bronchiolitis (ICD-9 466.XX) before 2nd birthday

**Exposure**  Residential exposure to primary fine particulate matter (PM2.5), nitrogen oxides (NOx), and carbon monoxide (CO) from traffic

**Control(s)**  Neighborhood socioeconomic status (SES), city region, child ethnicity, child sex, maternal asthma, Maternal smoking, birth year, and maternal age.

**Mediator(s)**  -

**Moderator(s)**  -

**Excluded/changed**

**Diagrams**

---

*Implied graph*

---

*DAG*
**Implied graph**

![Implied graph diagram](image)

28 edges assessed; 5 edges reversed; 12 edges retained; 11 edges removed

**Publication details #588**

<table>
<thead>
<tr>
<th>Author</th>
<th>Olivia Laugier, Patricia Garcia, Mohamed Boucékine et al.</th>
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</thead>
<tbody>
<tr>
<td>Year</td>
<td>2017</td>
</tr>
<tr>
<td><strong>Study title</strong></td>
<td>Influence of Socioeconomic Context on the Rehospitalization Rates of Infants Born Preterm</td>
</tr>
<tr>
<td><strong>Design</strong></td>
<td>Cohort study</td>
</tr>
<tr>
<td><strong>Methods</strong></td>
<td>Multivariable logistic regression analysis</td>
</tr>
<tr>
<td><strong>Sample</strong></td>
<td>3149 preterm infants with GA of &lt;32 + 6 weeks with residency in the West-Provence-Alpes-Côte-d'Azur region (France) and survived to the neonatal period</td>
</tr>
<tr>
<td><strong>Variables</strong></td>
<td></td>
</tr>
<tr>
<td><strong>Outcome</strong></td>
<td>Area-level deprivation</td>
</tr>
<tr>
<td><strong>Exposure(s)</strong></td>
<td>Emergency hospitalisation in first year of life (including bronchiolitis)</td>
</tr>
<tr>
<td><strong>Control(s)</strong></td>
<td>Gestational age, birth weight, SGA, sex, parity, bronchopulmonary dysplasia, urbanicity</td>
</tr>
<tr>
<td><strong>Mediator(s)</strong></td>
<td>-</td>
</tr>
<tr>
<td><strong>Moderator(s)</strong></td>
<td>-</td>
</tr>
<tr>
<td><strong>Excluded</strong></td>
<td>Necrotizing enterocolitis (inflammation of the bowel) and cerebral injury - related to other outcomes used in the this study</td>
</tr>
</tbody>
</table>

**DAG**

![DAG diagram](image)
**Implied graph**

![Diagram](image)

29 edges assessed; 13 edges reversed; 13 edges retained; 3 edges removed

### Publication details #6

| **Author** | Cintia Muñoz-Quiles, Mónica López-Lacort, Isabel Ubeda-Sansano et al. |
| **Year** | 2016 |
| **Study title** | Population-based Analysis of Bronchiolitis Epidemiology in Valencia, Spain |
| **Design** | Retrospective cohort; population and health databases in Valencia |
| **Methods** | Bayesian mixed Poisson regressions adjusted by fixed effects |
| **Sample** | 198,223 children |
| **Outcome** | Bronchiolitis hospitalisation <2 years |
| **Exposure(s)** | Social exclusion risk: obtained from electronic database (SIP), and its classification was based on multiple aspects such as unemployment, foreigner in irregular situation or without resources. |
| **Control(s)** | age, sex, year, prematurity, urban/rural residence (random effect: the month of the year (autocorrelated) and the health department) |
| **Mediator(s)** | - |
| **Moderator(s)** | - |
| **Excluded** | - |

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### Publication details #7

| **Author** | Khadra A Jama-Alol, Hannah C Moore, Peter Jacoby, Carol Bower and Deborah Lehmann |
| **Year** | 2014 |
| **Study title** | Morbidity due to acute lower respiratory infection in children with birth defects: a total population-based linked data study |
| **Design** | Retrospective cohort study |
**Methods**  | Negative binomial regression  
**Sample**  | 245,249 singleton births in Western Australia (1996-2005)  
**Variables**  
**Outcome**  | Number of ALRI hospitalisation before age 2 years (including Bronchiolitis <12 months)  
**Exposure(s)**  | Congenital anomalies  
**Control(s)**  | SEIFA index of disadvantage, sex, gestational age, birthweight, parity, season of birth, delivery method, Smoking during pregnancy, Asthma during pregnancy, maternal age, urban/rural  
**Mediator(s)**  | -  
**Moderator(s)**  | -  
**Excluded/changes**  | Birth defect changed to congenital anomalies  
**Diagrams**  
**Implied graph**  
![Implied graph image]  
**DAG**  
![DAG image]  
38 edges assessed; 18 edges reversed; 7 edges retained; 13 edges removed  
**Publication details**  
**Author**  | Hannah C Moore, Nicholas de Klerk, Patrick Holt, et al  
**Year**  | 2011  
**Study title**  | Hospitalisation for bronchiolitis in infants is more common after elective caesarean delivery  
**Design**  | Retrospective population-based data linkage cohort study, Western Australia  
**Methods**  | Negative binomial regression  
**Sample**  | 212 068 non-Aboriginal singleton births of 37–42 weeks gestation.
### Variables

<table>
<thead>
<tr>
<th><strong>Outcome</strong></th>
<th>Hospitalisations for bronchiolitis in children aged &lt;12 months</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Exposure(s)</strong></td>
<td>Delivery method</td>
</tr>
<tr>
<td><strong>Control(s)</strong></td>
<td>Socioeconomic index, pre-eclampsia, gestational diabetes, breech presentation, gestational age, smoking during pregnancy, maternal asthma, infant sex, season of birth, SGA, number of previous pregnancies, maternal age (years), year of birth,</td>
</tr>
<tr>
<td><strong>Mediator(s)</strong></td>
<td>-</td>
</tr>
<tr>
<td><strong>Moderator(s)</strong></td>
<td>-</td>
</tr>
<tr>
<td><strong>Excluded/changed</strong></td>
<td>Excluded number of admissions aged &lt;12 months. Reason: not applicable to bronchiolitis in infancy outcome. Proportion of optimal birth weight (POBW), a measure which takes into account gestational duration, fetal sex, maternal age, maternal height and parity: changed to SGA</td>
</tr>
</tbody>
</table>

### Diagrams

#### Implied graph *

![DAG](image1)

51 edges assessed; 11 edges reversed; 35 edges retained; 5 edges removed

#### DAG *

![DAG](image2)

### Publication details #9

<table>
<thead>
<tr>
<th><strong>Author</strong></th>
<th>Carme Puig, Jordi Sunyer, Oscar Garcia-Algar, et al.</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Year</strong></td>
<td>2008</td>
</tr>
<tr>
<td><strong>Study title</strong></td>
<td>Incidence and risk factors of lower respiratory tract illnesses during infancy in a Mediterranean birth cohort</td>
</tr>
<tr>
<td>-----------------</td>
<td>------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td><strong>Design</strong></td>
<td>Birth cohort recruited in Barcelona, Spain: home visit and follow-up questionnaire</td>
</tr>
<tr>
<td><strong>Methods</strong></td>
<td>Multivariate regression</td>
</tr>
<tr>
<td><strong>Sample</strong></td>
<td>A total of 487 infants were recruited at birth and followed up for 1 year</td>
</tr>
<tr>
<td><strong>Variables</strong></td>
<td></td>
</tr>
<tr>
<td><strong>Outcome</strong></td>
<td>LRTI in infancy including bronchiolitis</td>
</tr>
<tr>
<td><strong>Exposure(s)</strong></td>
<td>Paternal occupation using the UK Registrar General's 1990 classification</td>
</tr>
<tr>
<td><strong>Control(s)</strong></td>
<td>Maternal asthma, number of siblings, breastfed duration</td>
</tr>
<tr>
<td><strong>Mediator(s)</strong></td>
<td>-</td>
</tr>
<tr>
<td><strong>Moderator(s)</strong></td>
<td>-</td>
</tr>
<tr>
<td><strong>Excluded</strong></td>
<td>-</td>
</tr>
</tbody>
</table>

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*In the diagrams, the yellow node is the exposure, the red nodes are ancestors of the exposure and outcome, the blue node with a black outline is the outcome and all other blue nodes are ancestors of the outcome. Red edges represent biasing paths and green edges represent causal paths.*
Appendix 6.2. Lancet abstract: Could socioeconomic inequity in bronchiolitis admissions be reduced through intervening on gestational age?

Could socioeconomic inequity in bronchiolitis admissions be reduced through intervening on gestational age?  
A principled analysis of cohort data

Katie Marie Lewis, Bianca De Stavola, Fia Herbstfeld
Appendix 6.3. Stata code for causal mediation analyses

Set up file

******************************************************************************
* Sets up file for:
* 1) IPW of MSMs – for categorical X
* 2) G-estimation of structural nested model - for categorical X
* M-Y confounders c1 c2 c3 c4 c5 c6
* X-Y counfounders c1 c2 c6
******************************************************************************

use "$temp\cohort_main.dta", clear

* IPW of MSMs

gen id= childid

* Model X
mlogit x, rrr
predict num_1 num_2 num_3

*************

1) IPW of MSMs method

***firstly calculate baseline risk using IPW without BSs ***
**Step 1. model X
mlogit x, rrr
predict num_1 num_2 num_3
mlogit x i.c1 c2 i.c6, rrr
predict den_1 den_2 den_3

gen ipw1 = num_1/den_1 if x==1
replace ipw1 = num_2/den_2 if x==2
replace ipw1 = num_3/den_3 if x==3

su ipw1
cap drop num* den*

*model for Y (weighted) giving risk differences
binreg y i.x [pw=ipw],rd

// _b[_cons] = baseline risk

*************** run program for other estimates ***********************
cap program drop CDM_ipw
program CDM_ipw, rclass
version 15
args y x m c1 c2 c3 c4 c5 c6

cap drop num* den*
cap drop ipw*
cap drop X_M*

**Step 1. model X
mlogit `x', rrr
predict num_1 num_2 num_3

mlogit `x' i.`c1' `c2' i.`c6'; rrr
predict den_1 den_2 den_3

su num* den*

gen ipw1 = num_1/den_1 if `x'==1
replace ipw1 = num_2/den_2 if `x'==2
replace ipw1 = num_3/den_3 if `x'==3

su ipw1
cap drop num* den*

*************************************************************************
* Marginal risk difference using IPW
*************************************************************************
*** results
local mrd_2=_b[2.`x']
scalar mrd_2=`mrd_2'
return scalar mrd_2=`mrd_2'

local mrd_3=_b[3.`x']
scalar mrd_3=`mrd_3'
return scalar mrd_3=`mrd_3'

*** back to CDM ***

*Step 2. Model M
logit `m', or
predict num
logit `m' `x' i.`c1' `c2' `c3' i.`c4' `c5' i.`c6', or
predict den

su num den
gen ipw2 = cond(`m', num/den, (1-num)/(1-den))
su ipw2
cap drop num den

gen ipw=ipw1*ipw2
su ipw

*Step 3. model for Y (weighted) giving risk differences
gen X_M_2=(`x'==2)*`m'
gen X_M_3=(`x'==3)*`m'
binreg `y' i.`x' `m' X_M_2 X_M_3 [pw=ipw],rd

*** results CDM
local cdm_2=_b[2.`x']
scalar cdm_2=`cdm_2'
return scalar cdm_2=`cdm_2'

local cdm_3=_b[3.`x']
scalar cdm_3=`cdm_3'
return scalar cdm_3=`cdm_3'

*** results %
local per_2=(`mrd_2'-`cdm_2')/`mrd_2'*100
scalar per_2=`per_2'
return scalar per_2=`per_2'

local per_3=(`mrd_3'-`cdm_3')/`mrd_3'*100
scalar per_3=`per_3'
return scalar per_3=`per_3'
*check that it works:
CDM_ipw y x m i.c1 c2 c3 i.c4 c5 i.c6

*run with 1000 bootstraps
bootstrap MRD_2=r(mrd_2) MRD_3=r(mrd_3) ///
CDM_2=r(cdm_2) CDM_3=r(cdm_3) ///
PER_2=r(per_2) PER_3= r(per_3), reps(1000) seed(1234): ///
CDM_ipw y x m i.c1 c2 c3 i.c4 c5 i.c6
ex

2) **G-estimation of structural nested models method**

```stata
cap program drop CDM_snrm
program CDM_snrm, rclass
    version 15
    args y x m c1 c2 c3 c4 c5 c6
    cap drop m_hat
    cap drop r_M
    cap drop X_r_M
    cap drop X2_r_M X3_r_M
    cap drop Y_tilda2
    cap drop x_hat
    cap drop r_X
    cap drop r_X2 r_X3
    cap drop y1 y2 y3
    cap drop x1_hat x2_hat x3_hat

* Marginal risk difference
```

```stata
logit `y' i.`c1' `c2' i.`c6' if `x'=1
predict y1

logit `y' i.`c1' `c2' i.`c6' if `x'=2
predict y2

logit `y' i.`c1' `c2' i.`c6' if `x'=3
predict y3

sum y1, meanonly
local meany1=r(mean)
```
sum y2, meanonly
local meany2=r(mean)
sum y3, meanonly
local meany3=r(mean)
local risk1=`meany1'
local mrd2=`meany2' - `meany1'
local mrd3=`meany3' - `meany1'
scalar risk1=`risk1'
scalar mrd2=`mrd2'
scalar mrd3=`mrd3'
return scalar risk1=`risk1'
return scalar mrd2=`mrd2'
return scalar mrd3=`mrd3'
scalar list risk1 mrd2 mrd3

*** back to CDM ***

**Step 1a. predict M probabilities
logit `m' i.`x' i.`c1' `c2' `c3' i.`c4' `c5' i.`c6', or
predict m_hat

*Step 1b. Create M residuals
gen r_M=`m'-m_hat

**Step 2. model for Y
gen X2_r_M=(`x'==2)*r_M
gen X3_r_M=(`x'==3)*r_M
regress `y' r_M X2_r_M X3_r_M i.`c1' `c2' `c3' i.`c4' `c5' i.`c6'
scalar gamma2=_b[r_M]
scalar gamma3_2=_b[X2_r_M]
scalar gamma3_3=_b[X3_r_M]

**Step 3. intermediate transformation
gen Y_tilda2=`y'-gamma2*`m'-gamma3_2*(`x'==2)*`m'-gamma3_3*(`x'==3)*`m'

**Step 4a. predict X probabilities
mlogit `x' i.`c1' `c2' i.`c6', rrr
predict x1_hat x2_hat x3_hat

*Step 4b. Create X residuals
gen r_X2=(`x'==2)-x2_hat
gen r_X3=(`x'==3)-x3_hat

**Step 5. Regress transformed outcome
regress Y_tilda2 r_X2 r_X3 i.`c1' `c2' i.`c6'

*** results CDM
local
cdm2=_b[r_X2]
local
cdm3=_b[r_X3]
scalar
cdm2=`cdm2'
scalar
cdm3=`cdm3'
return
scalar cdm2=`cdm2'
return
scalar cdm3=`cdm3'
scalar
list cdm2 cdm3

*** results %
local per2=(`mrd2'-'cdm2')/'mrd2'*100
scalar per2=`per2'
return scalar per2=`per2'

local per3=(`mrd3'-'cdm3')/'mrd3'*100
scalar per3=`per3'
return scalar per3=`per3'

end

******************************************************************************
*check that it works:
CDM_snm y x m i.c1 c2 c3 i.c4 c5 i.c6

*run with 1000 bootstraps
bootstrap RISK_1= r(risk1) MRD_2=r(mrd2) MRD_3=r(mrd3) ///
          CDM_2=r(cdm2) CDM_3=r(cdm3) ///
          PER_2=r(per2) PER_3=r(per3), reps(1000) seed(1234): ///
          CDM_snmx y x m i.c1 c2 c3 i.c4 c5 i.c6
ex
******************************************************************************
## Appendix 7.1. Defining chronic conditions

**Table A7.1. ICD-10 codes defining chronic conditions (from Kristensen)**

<table>
<thead>
<tr>
<th>Group</th>
<th>ICD-10 code</th>
<th>Condition/s</th>
</tr>
</thead>
<tbody>
<tr>
<td>Malformations of the respiratory system</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Q30.0–Q30.9</td>
<td></td>
<td>Other malformations of the respiratory system</td>
</tr>
<tr>
<td>Q31.0–Q31.9</td>
<td></td>
<td>Malformations of the larynx; trachea and bronchi</td>
</tr>
<tr>
<td>Q32.0–Q32.4</td>
<td></td>
<td>Maleformations of the lungs</td>
</tr>
<tr>
<td>Q33.0–Q33.9</td>
<td></td>
<td>Maleformations of the lungs</td>
</tr>
<tr>
<td>Q34.0–Q34.9</td>
<td></td>
<td>Maleformations of the lungs</td>
</tr>
<tr>
<td>Q35.0–Q37.9</td>
<td></td>
<td>Cleft lip and palate</td>
</tr>
<tr>
<td>Other conditions associated with respiratory symptom</td>
<td></td>
<td></td>
</tr>
<tr>
<td>E84.0–E84.9</td>
<td></td>
<td>Cystic fibrosis</td>
</tr>
<tr>
<td>P27.0–P27.9</td>
<td></td>
<td>Bronchopulmonary dysplasia</td>
</tr>
<tr>
<td>Q39.0–Q39.1</td>
<td></td>
<td>Esophageal atresia</td>
</tr>
<tr>
<td>Q39.2–Q39.9</td>
<td></td>
<td>Other malformations of the esophagus</td>
</tr>
<tr>
<td>Q79.0</td>
<td></td>
<td>Congenital diaphragmatic hernia</td>
</tr>
<tr>
<td>Neuromuscular disease</td>
<td></td>
<td></td>
</tr>
<tr>
<td>G60.0–G60.9</td>
<td></td>
<td>Cerebral palsy</td>
</tr>
<tr>
<td>G70.2</td>
<td></td>
<td></td>
</tr>
<tr>
<td>G80.0–G80.9</td>
<td></td>
<td>Cerebral palsy</td>
</tr>
<tr>
<td>P94.1–P94.9</td>
<td></td>
<td>Congenital disturbances of muscle tonus, peripheral nerve disease, congenital myasthenia</td>
</tr>
<tr>
<td>Q01.0–Q01.9</td>
<td></td>
<td>Encephalocele</td>
</tr>
<tr>
<td>Q02.9</td>
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<td>Microcephalus</td>
</tr>
<tr>
<td>Q03.0–Q03.9</td>
<td></td>
<td>Congenital hydrocephalus</td>
</tr>
<tr>
<td>Q04.0–Q04.9</td>
<td></td>
<td>Other cerebral malformations</td>
</tr>
<tr>
<td>Q05.0–Q05.9</td>
<td></td>
<td>Spina bifida and malformations of the spinal cord</td>
</tr>
<tr>
<td>Q06.0–Q06.9</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Q07.0–Q07.9</td>
<td></td>
<td>Other malformations in the nervous system</td>
</tr>
<tr>
<td>G12.0–G12.9</td>
<td></td>
<td>Spinal muscular atrophy</td>
</tr>
<tr>
<td>G71.0–G71.3</td>
<td></td>
<td>Muscular dystrophy</td>
</tr>
<tr>
<td>Congenital diseases of the heart and the urinary system, chromosomal abnormalities, and others</td>
<td></td>
<td></td>
</tr>
<tr>
<td>D80.0–D82.9</td>
<td></td>
<td>Congenital immunodeficiencies</td>
</tr>
<tr>
<td>E70.0–E73.0</td>
<td></td>
<td>Inborn errors of metabolism</td>
</tr>
<tr>
<td>E74.0–E83.9</td>
<td></td>
<td></td>
</tr>
<tr>
<td>I27.0</td>
<td></td>
<td></td>
</tr>
<tr>
<td>N07.0–N07.9</td>
<td></td>
<td>Conditions affecting renal function</td>
</tr>
<tr>
<td>N13.0–N13.9</td>
<td></td>
<td>Malformations of the urinary system</td>
</tr>
<tr>
<td>N25.0–N25.9</td>
<td></td>
<td></td>
</tr>
<tr>
<td>P35.0–P35.9</td>
<td></td>
<td>Congenital viral infection</td>
</tr>
<tr>
<td>Q20.0–Q26.9</td>
<td></td>
<td>Congenital heart disease</td>
</tr>
<tr>
<td>Code</td>
<td>Description</td>
<td></td>
</tr>
<tr>
<td>----------</td>
<td>-----------------------------------------------------------</td>
<td></td>
</tr>
<tr>
<td>Q40.0–Q43.9</td>
<td>Malformations of the gastrointestinal tract, liver, biliary system, pancreas, and the abdominal wall</td>
<td></td>
</tr>
<tr>
<td>Q44.0–Q44.7</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Q45.0–Q45.3</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Q60.0–Q64.9</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Q79.2–Q79.3</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Q86.0</td>
<td>Fetal alcohol syndrome</td>
<td></td>
</tr>
<tr>
<td>Q90.0–Q90.9</td>
<td>Down syndrome</td>
<td></td>
</tr>
<tr>
<td>Q91.0–Q99.9</td>
<td>Other chromosomal abnormalities</td>
<td></td>
</tr>
</tbody>
</table>
Appendix 7.2. Example Mplus code

**Deriving asthma/wheeze trajectory groups**

! CODE FOR LCGA 4 CLASSES CUBIC MODEL
! TERMS TO INCLUDE FOR GMM ALSO SHOWN
! MANUAL 3 STEP APPROACH

TITLE: Y on LCGA 4 CLASS ASTHMA/WHEEZE: AW2-AW9

!AWx = asthma/wheeze at age x

DATA: FILE IS "file_name.dat";
    Format is free;

VARIABLES:
    NAMES = AW2-AW9 ID ;
    USEV = AW2-AW9;
    MISSING are all (-9999);
    IDVARIABLE = CHILDID;
    CLASSES = C(4)
    CATEGORICAL = AW2-AW9;

ANALYSIS:
    TYPE = MIXTURE;
    ALGORITHM = INTEGRATION;
    STARTS = 500 100;
    STITERATIONS = 10;
    PROCESS = 10(STARTS);

MODEL:
! This describes the model to be estimated
! is q cu are names for intercept, slope, quadratic and cubic terms, respectively

    %overall%
    i s q cu | aw2@0 aw3@1 aw4@2 aw5@3 aw6@4 aw7@5 aw8@6 aw9@7;
    i-cu @ 0;

! Delete "@ 0" to allow within class variation (GMM model)
! This allows for the additional terms needed for GMM models

OUTPUT : TECH1 TECH8 TECH11 TECH 14 CINTERVAL;

! TECH1 produces starting values for all free parameters in the model

*Add details from best fitting model into the 11 imputed datasets*
! CODE FOR adding LCGA details into imputation datasets
! Repeat on each imputed dataset
! use parameters from output of above code to replicate the same LCGA output in each
! dataset

TITLE: Adding LCGA 4 CLASS ASTHMA/WHEEZE

DATA: FILE IS "imputed_dataset_1.dat";
Format is free;

VARIABLES:
   NAMES = AW2-AW9 ID SGA area_groups asthma_mother season birthyear bronch chronic_cond delivery_meth deprivation3 mat_age mother_born _uk preterm sex siblings smoke_preg;
   USEV = AW2-AW9;
   MISSING are all (-9999);
   IDVARIABLE = ID;
   CLASSES = C(4)
   CATEGORICAL = AW2-AW9;
   AUXILIARY = SGA area_groups asthma_mother season birthyear bronch chronic_cond delivery_meth deprivation3 mat_age mother_born _uk preterm sex siblings smoke_preg;

ANALYSIS:
   TYPE = MIXTURE;
   ALGORITHM = INTEGRATION;
   STARTS = 0;
   !OPTSEED = 2341;

MODEL:
   %overall%
   i s q cu | aw2@0 aw3@1 aw4@2 aw5@3 aw6@4
               aw7@5 aw8@6 aw9@7;

! Class-specific means
   [c#1*3.931];
   [c#2*3.745];
   [c#3*3.584];

! Class-specific growth factors from the final LCGA model
   %C#1%
   [i*4.983];
   [s*1.635];
   [q*-0.269];
   [cu*0.010];
   i-cu@0;


```
\%C#2%
[i*1.380];
[s*0.855];
[q*0.079];
[cu*-0.016];
i-cu@0;

\%C#3%
[i*4.935];
[s*1.047];
[q*-0.446];
[cu*0.036];
i-cu@0;

\%C#4%
[i*0];
[s*-0.210];
[q*-0.058];
[cu*0.013];
i-cu@0;

OUTPUT : FILE is LCGA_step1_imp1.dat;
save is Cprob;

Multinomial logistic regression

! CODE FOR multinomial logistic regression looking at risk factors by trajectory group
! example of SIMD group as exposure
! uses all imputed datasets and Rubin’s rules applied

TITLE: Multinomial logistic regression

DATA: FILE IS
  "\imputed_dataset_list.dat";
    Format is free;
    Type = imputation;

VARIABLES:
  NAMES = AW2-AW9 SGA area_groups asthma_mother season
         birthyear bronch chronic_cond delivery_meth deprivation3 mat_age
         mother_born _uk preterm sex siblings smoke_preg Cprobs1-Cprobs4 N ID;

USEV = N SIMD1 SIMD2 SIMD3;

IDVARIABLE = ID;
CLASSES = C(4)
NOMINAL = N;
```
DEFINE:
    SIMD1=0;
    SIMD2=0;
    SIMD3=0;
    If (deprivation3 eq 3) then SIMD1=1;
    If (deprivation3 eq 2) then SIMD2=1
    If (deprivation3 eq 1) then SIMD3=1

ANALYSIS:
    TYPE = MIXTURE;
    STARTS = 0;

MODEL:
    %overall%
    C ON SIMD1 SIMD2 SIMD3;

! including logit classification probabilities from LCGA model
! read table output by row

    %C#1%
    [N#1@5.186];
    [N#2@2.229];
    [N#3@1.917];

    %C#2%
    [N#1@-1.113];
    [N#2@2.068];
    [N#3@-0.542];

    %C#3%
    [N#1@-1.520];
    [N#2@-1.546];
    [N#3@1.525];

    %C#4%
    [N#1@-13.815];
    [N#2@8.62];
    [N#3@-7.754];

OUTPUT : CINTERVAL;
Appendix 7.3. Stata code for mediation analysis

Set up file and run analyses in one

************************************************
* File for:
* 1) overall % chronic asthma attributable to bronchiolitis
* 2) IPW of MSMS – for categorical X
* begins with imputation
*X: deprivation3
*M: bronch
*Y: binary_asthma
*M-Y confounders: area_groups, birthyear_child, mother_born_UK
*X-Y confounders: area_groups, birthyear_child, mother_born_UK, birth_season, sex,
   mat_age_cat, preterm, smoke_preg, chronic_conditions, SGA, asthma_mother, siblings
*keep birthweight, gestation, firstborn to calculate SGA post imputation
*also keep lbw, mat_age, socio_economic_group, intensive_flag, preterm_conditions,
hospital_code
************************************************
use     "$temp\CDMasthma_dataset.dta", clear
rename deprivation3 x
rename bronch m
generate y=0
replace y=1 if class==1 | class==2
rename area_groups c1
gen     c2=0
replace c2=1 if birthyear_child==2008
rename mother_born_UK c3
rename birth_season c4
rename sex c5
rename mat_age_cat c6 // not in imputation model
rename preterm c7 // not in imputation model
rename smoke_preg c8
rename chronic_condition c9
rename SGA c10 // not in imputation model
rename asthma_mother c11
rename siblings_cat c12 // not in imputation model
rename delivery_method_3cat c13
replace c13=c13-1 //so categorised 0 1 2
rename preterm_conditions iv1
rename birthweight iv2
rename gestation iv3
rename mat_age iv4
rename intensive_flag iv5
rename socio_economic_group iv6
rename hosp_G iv7
rename siblings iv8

************************************************************************
cap program drop CDM_asthma
program define CDM_asthma, rclass

******************************************************
*SINGLE IMPUTATION
*for missing data

*FOR iv2 = birthweight (to create SGA c10) regress
*FOR iv3 = gestation (to create preterm c7) regress
*FOR iv4 = mat_age (to create mat age cats c13) regress
*FOR iv8 = siblings (to create categorical siblings c12 and SGA c10 too) mlogit
*FOR c13 = delivery_method mlogit
*FOR c8 = smoke_preg logit
*FOR c11 = asthma_mother logit

************************************************************************
***generate imputation vars
qui foreach var in x m y c1 c2 c3 c4 c5 c6 c7 ///
    c8 c9 c10 c11 c12 c13 ///
    iv1 iv2 iv3 iv4 iv5 iv6 iv7 iv8 {
    gen imp_`var'=`var'
}

***replace missingness in vars with average values
*binary vars with missing values
qui foreach var in c8 c11 {
    summ `var'
    replace imp_`var'=runiform()<r(mean) if imp_`var'==.
}

*continuous vars with missing values
qui foreach var in iv2 iv3 iv4 iv8 {
    summ `var'
    replace imp_`var'=r(mean) if imp_`var'==.
}

*categorical var with missing values
qui ta c13,m
qui gen imp_c13_0=(c13==0)
qui gen imp_c13_2=(c13==2)
qui replace imp_c13_0=. if c13==.
qui replace imp_c13_2=. if c13==.
qui replace imp_c13_0=runiform()<0.05 if imp_c13_0==.
qui replace imp_c13_2=0 if imp_c13_0==1
qui replace imp_c13_2=runiform()>0.81 if imp_c13_2==.
qui replace imp_c13=imp_c13_0+imp_c13_2*2 if imp_c13==.

qui su imp_*

*Need to do a lead in to imputation to stop convergence
forvalues cycle=1(1)10 {

***impute binary vars first
*for smoke_preg c8
qui logit c8 i.imp_x imp_m imp_y i.imp_c1 imp_c3 i.imp_c4 ///
    imp_c5 imp_c9 imp_c11 i.imp_c13 ///
    imp_iv1 imp_iv2 imp_iv3 imp_iv4 imp_iv5 imp_iv6 imp_iv7 imp_iv8
qui predict ec8
qui replace imp_c8=runiform()<ec8 if c8==.
qui tab imp_c8
qui drop ec8

*for asthma_mother c11
qui logit c11 i.imp_x imp_m imp_y i.imp_c1 imp_c3 i.imp_c4 imp_c5 ///
    imp_c8 imp_c9 i.imp_c13 ///
    imp_iv1 imp_iv2 imp_iv3 imp_iv4 imp_iv5 imp_iv6 imp_iv7 imp_iv8
qui predict ec11
qui replace imp_c11=runiform()<ec11 if c11==.
qui tab imp_c11
qui drop ec11

***impute continuous vars
*for birthweight iv2
qui regress iv2 i.imp_x imp_m imp_y i.imp_c1 imp_c3 i.imp_c4 imp_c5 ///
    imp_c8 imp_c9 imp_c11 i.imp_c13 ///
    imp_iv1 imp_iv3 imp_iv4 imp_iv5 imp_iv6 imp_iv7 imp_iv8
qui predict eiv2
qui replace imp_iv2=eiv2+e(rmse)*rnormal() if iv2==.
qui drop eiv2

*for gestation iv3
qui regress iv3 i.imp_x imp_m imp_y i.imp_c1 imp_c3 i.imp_c4 imp_c5 ///
    imp_c8 imp_c9 imp_c11 i.imp_c13 ///
    imp_iv1 imp_iv2 imp_iv4 imp_iv5 imp_iv6 imp_iv7 imp_iv8
qui predict eiv3
qui replace imp_iv3=eiv3+e(rmse)*rnormal() if iv2==.
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qui drop eiv3

*for mat_age iv4
qui regress iv4 i.imp_x imp_m imp_y i.imp_c1 imp_c3 i.imp_c4 imp_c5 ///
    imp_c8 imp_c9 imp_c11 i.imp_c13 ///
    imp_iv1 imp_iv2 imp_iv3 imp_iv5 imp_iv6 imp_iv7 imp_iv8
qui predict eiv4
qui replace imp_iv4=eiv4+e(rmse)*rnormal() if iv4==.
qui drop eiv4

*for siblings iv8
qui regress iv8 i.imp_x imp_m imp_y i.imp_c1 imp_c3 i.imp_c4 imp_c5 ///
    imp_c8 imp_c9 imp_c11 i.imp_c13 ///
    imp_iv1 imp_iv2 imp_iv3 imp_iv4 imp_iv5 imp.iv6 imp.iv7
qui predict eiv8
qui replace imp_iv8=eiv8+e(rmse)*rnormal() if iv8==.
qui drop eiv8

***impute categorical var
*for delivery_method c13
qui mlogit c13 i.imp_x imp_m imp_y i.imp_c1 imp_c3 i.imp_c4 imp_c5 ///
    imp_c8 imp_c9 imp_c11 ///
    imp_iv1 imp_iv2 imp_iv3 imp_iv4 imp_iv5 imp_iv6 imp_iv7 imp_iv8
qui predict ec13_0 ec13_1 ec13_2
qui replace imp_c13_0=runiform()<ec13_0 if c13==.
qui replace imp_c13_2=0 if imp_c13_0==1
qui replace imp_c13_2=runiform()>1-ec13_2) if c13==.
qui replace imp_c13=(imp_c13_0==1)*0+(imp_c13_0==0)*(imp_c13_2==0)*1 ///
    + (imp_c13_2==1)*2 if c13==.
qui tab imp_c13
qui drop ec13*
}

*keep orginal vars only
qui foreach var of varlist iv2 iv3 iv4 iv8 c8 c11 c13 {
    replace `var'=imp_`var' if `var'==.
}
qui drop imp_*

***create the new vars needed for analysis
*preterm c7 from gestation iv3
qui replace c7=0 if iv3<37 & c7==.
qui replace c7=1 if iv3>=37 & c7==.

*mat age cats c6 from mat_age iv4
qui replace c6=0 if iv4<=19 & c6==.
qui replace c6=1 if (iv4>19 & iv4<30) & c6==.
qui replace c6=2 if (iv4>29 & iv4<40) & c6==.
qui replace c6=3 if iv4>=40 & c6==.

*categorical siblings c12 from siblings iv8
qui replace c12=0 if (round(iv8)==0 | iv8<0) & c12==.
qui replace c12=1 if round(iv8)==1 & c12==.
qui replace c12=2 if round(iv8)>1 & c12==.

*SGA c10 from birthweight iv2 gestation iv3 and siblings c12
qui generate firstborn=1 if c12==0
qui replace firstborn=0 if c12==1 | c12==2
qui generate gestation=round(iv3)
qui replace gestation=24 if gestation==23
qui replace gestation=43 if gestation>43
qui generate sex=c5
qui merge m:m sex gestation using "$temp\SGA.dta"
qui replace c10=0 if iv2>tenth_centile & c10==.
qui replace c10=0 if iv2<=tenth_centile & c10==.
qui drop _merge tenth_centile gestation firstborn sex

************************************************************************
* CDM analysis
**************************************************
********************
cap drop num*
cap drop den*
cap drop ipw*
cap drop X_M*

*1. Model x
qui mlogit x, base(1) rrr
qui predict num_1 num_2 num_3
qui mlogit x i.c1 c2 c3, base(1) rrr
qui predict den_1 den_2 den_3
qui gen ipw1 = num_1/den_1 if x==1
qui replace ipw1 = num_2/den_2 if x==2
qui replace ipw1 = num_3/den_3 if x==3

*model for asthma_binary (weighted) giving risk differences for TCE
qui binreg y i.x [pw=ipw],rd
local tce_2=_b[2.x]
scalar tce_2="tce_2"
return scalar tce_2='tce_2'
local tce_3=_b[3.x]
scalar tce_3=`tce_3'
return scalar tce_3=`tce_3'

qui cap drop num* den*

*Step 2. Model M
qui logit m, or
qui predict num
qui logit m i.x i.c1 c2 c3 i.c4 c5 i.c6 c7 c8 c9 c10 i.c12 i.c13
qui predict den
qui gen ipw2 = cond(m, num/den, (1-num)/(1-den))
qui cap drop num* den*
qui cap drop ipw
qui gen ipw=ipw1*ipw2

*Step 3. model for Y (weighted) giving risk differences
qui gen X_M_2=(x==2)*m
qui gen X_M_3=(x==3)*m
qui binreg y i.x X_M_2 X_M_3 [pw=ipw],rd

*** Counterfactual Disparity Measures
local cdm_2=_b[2.x]
scalar cdm_2=`cdm_2'
return scalar cdm_2=`cdm_2'

local cdm_3=_b[3.x]
scalar cdm_3=`cdm_3'
return scalar cdm_3=`cdm_3'

local risk_1=_b[_cons]
scalar risk_1=`risk_1'
return scalar risk_1=`risk_1'

*** results %
local per_2=(`tce_2'-'cdm_2')/tce_2'*100
scalar per_2=`per_2'
return scalar per_2=`per_2'

local per_3=(`tce_3'-'cdm_3')/tce_3'*100
scalar per_3=`per_3'
return scalar per_3=`per_3'

end
*check that it works:
CDM_\text{asthma} \quad y \times m \ \text{i.c1} \ \text{c2} \ \text{i.c3} \ \text{c4} \ \text{i.c5} \ \text{c6} \ \text{c7} \ \text{c8} \ \text{c9} \ \text{c10} \ \text{i.c11} \ \text{i.c12} \ \text{i.c13}

*Bootstrap
bootstrap 
TCE_2=r(tce_2) \ \text{TCE}_3=r(tce_3) \ \text{RISK}_1=r(risk_1) \\
CDM_2=r(cdm_2) \ \text{CDM}_3=r(cdm_3) \\
\text{PER}_2=r(per_2) \ \text{PER}_3=r(per_3) \\
, \ \text{reps}(1000) \ \text{seed}(1234): \ \text{CDM}_\text{asthma}
### Appendix 7.4. Risk factor distribution by asthma/wheeze trajectories

#### Table A7.2. Risk factor distribution by asthma/wheeze trajectories (groups based on most likely class membership)

<table>
<thead>
<tr>
<th>Factor</th>
<th>Never or infrequent (N=78,803)</th>
<th>Early-Transient (N=1,853)</th>
<th>Early-Persistent (N=1,547)</th>
<th>Intermediate-onset (N=1,645)</th>
</tr>
</thead>
<tbody>
<tr>
<td>SIMD</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Low</td>
<td>26.6%</td>
<td>20.5%</td>
<td>19.8%</td>
<td>21.4%</td>
</tr>
<tr>
<td>Medium</td>
<td>37.9%</td>
<td>36.8%</td>
<td>37.0%</td>
<td>37.9%</td>
</tr>
<tr>
<td>High</td>
<td>35.5%</td>
<td>42.7%</td>
<td>43.2%</td>
<td>40.7%</td>
</tr>
<tr>
<td>≥1 bronchiolitis admission</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>No</td>
<td>97.2%</td>
<td>89.2%</td>
<td>91.0%</td>
<td>94.5%</td>
</tr>
<tr>
<td>Yes</td>
<td>2.8%</td>
<td>10.8%</td>
<td>9.0%</td>
<td>5.5%</td>
</tr>
<tr>
<td>Birth season</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Spring</td>
<td>33.4%</td>
<td>30.5%</td>
<td>34.1%</td>
<td>35.3%</td>
</tr>
<tr>
<td>Summer</td>
<td>22.8%</td>
<td>23.8%</td>
<td>23.7%</td>
<td>22.7%</td>
</tr>
<tr>
<td>Autumn</td>
<td>16.8%</td>
<td>17.1%</td>
<td>15.4%</td>
<td>15.9%</td>
</tr>
<tr>
<td>Winter</td>
<td>27.0%</td>
<td>28.7%</td>
<td>26.8%</td>
<td>26.1%</td>
</tr>
<tr>
<td>Sex</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Female</td>
<td>49.5%</td>
<td>39.2%</td>
<td>37.4%</td>
<td>40.2%</td>
</tr>
<tr>
<td>Male</td>
<td>50.5%</td>
<td>60.8%</td>
<td>62.6%</td>
<td>59.8%</td>
</tr>
<tr>
<td>Area</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Northern</td>
<td>13.0%</td>
<td>12.0%</td>
<td>12.7%</td>
<td>12.6%</td>
</tr>
<tr>
<td>Central</td>
<td>28.5%</td>
<td>28.4%</td>
<td>26.0%</td>
<td>25.5%</td>
</tr>
<tr>
<td>Glasgow</td>
<td>23.1%</td>
<td>25.4%</td>
<td>24.9%</td>
<td>26.1%</td>
</tr>
<tr>
<td>Edinburgh</td>
<td>17.1%</td>
<td>13.6%</td>
<td>14.1%</td>
<td>16.5%</td>
</tr>
<tr>
<td>Southern</td>
<td>18.2%</td>
<td>20.6%</td>
<td>22.4%</td>
<td>19.2%</td>
</tr>
<tr>
<td>Maternal birth country</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Non-UK</td>
<td>10.9%</td>
<td>9.1%</td>
<td>8.7%</td>
<td>8.9%</td>
</tr>
<tr>
<td>UK</td>
<td>89.1%</td>
<td>90.9%</td>
<td>91.3%</td>
<td>91.1%</td>
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<tr>
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