Tailoring Treatments using Treatment Effect Modification.

A F Schmidt\textsuperscript{a,b,c,d}, O H Klungel\textsuperscript{a,b}, M Nielen\textsuperscript{c}, A de Boer\textsuperscript{b}, R H H Groenwold\textsuperscript{a,b}, A W Hoes\textsuperscript{a}.

\textsuperscript{a} Julius Center for Health Sciences and Primary Care, University Medical Center Utrecht, P.O. Box 85500, 3508 GA Utrecht, the Netherlands.

\textsuperscript{b} Division of Pharmacoepidemiology and Clinical Pharmacology, Utrecht Institute for Pharmaceutical Sciences, P.O. Box 80082, 3508 TB Utrecht, The Netherlands.

\textsuperscript{c} Department of Farm Animal Health, Faculty of Veterinary Medicine, Utrecht University, Yalelaan 7, Utrecht 3584 CL, The Netherlands.

\textsuperscript{d} Institute of Cardiovascular Science, Faculty of Population Health, University College London, London WC1E 6BT, the United Kingdom.

* Contact: 0044 (0)20 3549 5625

E-mail address: amand.schmidt@ucl.ac.uk (A.F.Schmidt).

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None of the authors of this paper have a financial or personal relationship with other people or organisations that could inappropriately influence or bias the content of the paper.

Author contributions

AFS drafted the manuscript. OHK, MN, AB, AWH and RHHG provided guidance during initial planning of the paper and during critical revision.

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Prior postings and presentations

This study and its results have not been previously published, neither has it been presented at conferences.
Key Points

- Clinical studies are designed to provide evidence on average treatment effects.
- To tailor treatment towards individual patients, the presence or absence of treatment effect modification needs to be systematically elucidated.
- Generalizability of treatment effects can be tested within the framework of equivalence testing.
- The type of patient, the presence, the magnitude, and the number of effect modifiers determines whether no further analyses, univariable subgroup analyses, or multivariable subgroup analyses may need to be performed.
Abstract

Applying results from clinical studies to individual patients can be a difficult process. Using the concept of treatment effect modification (also referred to as interaction), defined as a difference in treatment response between patient groups, we discuss whether and how treatment effects can be tailored to better meet patients’ needs. First we argue that, contrary to how most studies are designed, treatment effect modification should be expected. Second, given this expected heterogeneity, a small number of clinically relevant subgroups should be a priori selected, depending on the expected magnitude of effect modification, and prevalence of the patient type. Third, by defining generalizability as the absence of treatment effect modification we show that generalizability can be evaluated within the usual statistical framework of equivalence testing. Fourth, when equivalence cannot be confirmed, we address the need for further analyses, and studies tailoring treatment towards groups of patients with similar response to treatment. Fifth, we argue that to properly frame, the entire body of evidence on effect modification should be quantified in a prior probability.
Background

Before launching a new treatment on the market, medical interventions and most notably drugs, are typically evaluated in randomized clinical trials (RCTs) which primarily focus on the intended effects of interventions. Sometimes, RCTs can also provide information on relatively common unintended (i.e. adverse) effects\textsuperscript{1-3}. After marketing, intervention effects (both intended and unintended) are often monitored using nonrandomized studies (e.g., case-control or cohort studies), supplemented by post-launch RCTs when needed. These studies are usually designed to provide information on the average intervention effect. Therefore, differences in treatment effects between a wide range of potential users will often remain undetected\textsuperscript{4-7}.

When treatment effects differ between patients, this is referred to as effect modification, interaction, or heterogeneity of treatment effects. Consider a hypothetical trial (Table 1) that includes patients with diabetes (40%) and patients without (60%). The risk ratio (RR) of the intervention effect on the 5-years incidence of stroke, differs between patients with and without diabetes: e.g. RR = 0.75 among patients with diabetes and RR = 0.63 among patients without diabetes. The observed (average) intervention effect is a weighted average of the effects among patients with and patients without diabetes: RR = 0.68. In this example, the intervention effect differs between subgroups based on diabetes status, i.e., there is effect modification by diabetes. Patients may be treated suboptimally when effect modification is not recognized.

Throughout this paper, we will use the term effect modification, interaction and heterogeneity interchangeably. Some reserve the term interaction for the specific
situation of heterogeneity of treatment effect when a factor biologically interacts with the
treatment and *effect modification* for the situation where it does not. This distinction can
usually not be determined analytically and will not be made here either. Also, it has been
recognized that the presence of effect modification depends on the effect measure
chosen. In the example RCT (table 1) there was interaction on the RR (and on the
risk difference [RD]) scale, however using the odds ratio (OR) the effect of treatment
was 0.25, in both diabetic and non-diabetic patients. Effect modification is therefore
also referred to as effect measure modification. Here, we consider situations where the
effect measure was selected a priori and thus only consider effect modification of the
particular effect measure chosen.

In this paper we build upon work by others, and use the concept of treatment effect
modification to discuss how generalizability of treatment effects can evaluated, and, if
generalizability is absent, how to tailor effects to patients with a more homogenous
treatment response.

**Should treatment effect modification be expected?**

Most clinical studies are not designed to detect treatment effect modification and usually
assume homogeneity of treatment effects. Because of this, power to detect interaction
effects is generally low, and absence of significant interaction effects should not be seen
as proof for the absence of treatment effect modification (a point we will revisit). Despite
this expected low power, Poole, Shrier and VanderWeele describe that between 34%
to 47% of the meta-analyses reviewed by Engels et.al., Deeks and Sterne and
Egger, rejected homogeneity of treatment effects. This, perhaps unexpected, high
percentage of heterogeneity is likely not solely attributable to differences in treatment response but may also be explained by between study differences in dosage, adherence strategies or co-medication (see Sun et.al.\textsuperscript{13;21} and Rothwell\textsuperscript{22}, for a more complete discussion).

Given the above mentioned problems of empirical evidence for treatment effect heterogeneity we refer to theoretical work of Greenland who showed that if both treatment and a potential effect modifier have an effect on the outcome, treatment effect modification must be present on at least one effect measure scale\textsuperscript{23} (e.g., RD or RR).

Given that most human diseases are complex in nature, multiple factors will be involved in a wide range of endpoints. Combining this with the tendency of more representative studies\textsuperscript{24;25}, and therefore more heterogeneous patient samples, we feel that some degree of treatment effect modification should be expected in most studies in which treatment has an effect on an outcome. Whether this effect modification is relevant for clinical practice, is a difficult question, which should be explored case by case.

**Which potential effect modifiers to pre-specify.**

An essential question when designing a study is for whom we want to assess the effects of treatment, whether treatment effects may differ, and if so what defines the subgroups for which treatment effects may differ. To pre-specify potential effect modifier it seems sensible to take account of any prior knowledge of the biological mechanism, potential patient benefit, the frequency certain patients are encountered in practice, and the costs involved in measuring a patient characteristic. When e.g. comorbidity is a potential effect modifier, it seems more reasonable to assess whether relatively common diseases, such
as diabetes, modify the effect of treatment. Discussions on the choice of subgroups should focus on patients included but also certainly on patients not included in a (future) study\textsuperscript{26}.

Too often, however, discussions on generalizability or the absence of treatment effect modification revolve around the question whether a patient sample is \textit{representative} of the target population or the “average” patient\textsuperscript{27}. Representativeness however, plays only a minor role in applying treatment effects to individuals\textsuperscript{14,25,28,29}. In the absence of effect modification the same treatment effect applies to every patient subgroup, and thus, representativeness is irrelevant. In the presence of treatment effect modification, due to unequal subgroup sizes, a representative sample will more often than not preclude detection of treatment effect modification. Hence, representativeness often results in wrongfully assuming homogeneity of treatment effects and thus possibly in patients being treated suboptimally. A more fruitful approach when expecting treatment effect modification is to design a study to oversample the pre-specified patient subgroups to ensure sufficient power to detect interaction or its absence.

Even if one is interested in population average treatment effect\textsuperscript{30} one should be aware that in the presence of treatment effect modification, small differences between populations can result in markedly different main treatment effects\textsuperscript{31}. Assume, for example that, in a population aged 65, the main treatment effect is 1.00 (RR). In the presence of an interaction effect of 0.95 (RR) per year, the treatment effect in a population aged 70 will be 0.77 (RR) \[i.e., e^{\ln(1.00)+\ln(0.95)\times(70-65)} \approx 0.77\]. Hence, unless
treatment effect modification is minimal, population average treatment effects are not expected to generalize to other settings.

Thus, when discussing generalizability or treatment effect modification, it is essential to define the patient group(s) of interest. Such subgroups should be chosen based on biological plausibility, potential patient benefit, subgroup frequency, and measurement costs, this should, however, not be guided by the issue of representativeness.

**When are treatment effects generalizable?**

When interaction effects can be quantified with sufficient precision to exclude clinically relevant treatment effect modification, the main (i.e. average) treatment effect equally applies to all subjects studied and - because there is no direct reason to believe the treatment acts differently in other subjects – this treatment effect is possibly generalizable to, and perhaps beyond, the population included in the study\(^{30;31}\). As stated previously non-significant interaction tests are not sufficient to claim generalizability; to quote Altman\(^{32}\) “absence of evidence is not evidence of absence”. Instead to ‘prove’ generalizability, so called equivalence tests should be used.

Recognizing that the strict null-hypothesis (i.e., \(H_0: \mu_0 = \text{null}\)) probably never holds, tests of equivalence determine margins between which differences in treatment effect estimates are small enough to be deemed clinically irrelevant\(^{33;34}\). When the treatment effect estimate and its confidence interval fall between these margins, equivalence is ‘proven’ (Figure 1). Equivalence tests can be applied to interactions effects by determining a margin around the neutral interaction effect or around the subgroup
specific effects, and testing if both the point estimates and their confidence intervals fall within this margin. For example, let $d$ be the predefined margin of equivalence, $\delta_i$ the effect for the $i$th subgroup, when $i = \{0,1\}$ the interaction effects equals $\theta = \delta_0 - \delta_1$, $\sigma_i$ the subgroup specific standard errors of $\delta_i$, and $\sigma_\theta$ the standard error of the interaction effect. In this case an interaction effect is sufficiently absent when $(\delta_i - z * \sigma_i, \delta_i + z * \sigma_i) \subset (-d, d)$ is true for all subgroup effects or $(\theta - z * \sigma_\theta, \theta + z * \sigma_\theta) \subset (-d, d)$, where $z = \phi^{-1}(1 - \frac{\alpha}{2k})$, with typically $\alpha = 0.05$ and $k$ equalling the number of subgroups or 1 if using an interaction effect. These procedures test against the following null-hypotheses

$H_0$: $|\delta_i| < d$ and $H_0$: $|\theta| < d$. The subgroup specific equivalence test is appealing because it requires complete equivalence in every subgroup, however as shown in Figure 2 power is likely low. Using the same margin as for the subgroup specific effects, an interaction equivalence test is more powerful (Figure 2). However, when using the interaction equivalence test with more than 2 underlying subgroups, some subgroup specific effects may violate the equivalence margins which may be undesirable. A clear benefit of the interaction equivalence tests over its subgroup specific counter parts is that it straightforwardly extends to linear effect modifiers (e.g., age), preventing arbitrary categorizations.

**Detecting treatment effect modification.**

Effect modification can be detected by testing whether the interaction effect differs from zero\textsuperscript{35,36}. However, such interaction tests are renowned for their lack of power (i.e., the probability of correctly concluding that an interaction exists) which may be compounded by large type 1 errors (i.e., the probability of falsely concluding that an interaction exists)
when the data are sparse\textsuperscript{10,37-43}. Note, data sparseness is intuitively defined as a small expected cell count but generalizes to continuous data with low densities (or frequencies) at certain values. Often this underperformance of interaction tests is viewed as inevitable; however, this underperformance is merely a result of a lack of a proper design to detect interaction effect; often resulting in sparse data.

For more definitive conclusions on the absence or presence of treatment effect modification, the current approach to interaction testing needs improvement. A first step is to more actively share and pool individual patient data to increase the effective sample size, power, and by decreasing data sparseness, ensuring nominal type 1 error rates for interaction tests\textsuperscript{31,44,45}.

Second, for interaction tests to be anything but exploratory, interaction tests should not only be pre-specified but also include proper sample size calculations and sampling strategies (e.g., equally sized subgroups); ensuring appropriate power and type 1 error rates. One attractive idea is to incorporate interaction tests using adaptive trial designs\textsuperscript{46-48}. For example, consider an RCT of a particular treatment, conducted within a homogenous group of patients. If during interim analysis there is enough evidence to expect that the treatment is effective (i.e. there is a beneficial average effect), the second study period (the period following the interim analysis) can be used to enrich the patient sample to explore heterogeneity between pre-specified clinically important patient subgroups. We recognize that this contrasts with the more usual approach of focusing on a single promising subgroup after interim\textsuperscript{46,49}. Here we actually reverse the usual approach; we start with a subgroup where we expect treatment to be most
beneficial and in the second stage (after interim) explore consistency of this treatment effect across important subgroups.

Third, to increase the interpretability of interaction tests (or any test) we feel it is essential to a priori define the prior probability of rejecting a test. For example let’s assume that data from multiple well designed studies are available, ensuring sufficient power (let’s say 70%, or type 2 error rate of 0.30) to reject an interaction test using a statistical significance level (alpha) of 0.05. Suppose two different drug compounds are evaluated, for the first compound we know that for a similar drug 1% of interaction test were true positives, for the second compound this was 25%. In the first case the probability that a rejected interaction test reflects a true positive equals 1 –

\[
1 - \frac{\alpha(1-prior)}{\alpha(1-prior) + prior(1-\beta)} = 1 - \frac{0.05(1-0.01)}{0.05(1-0.01) + 0.01(1-0.30)} = 0.13
\]

while for the second compound this equals 0.82. Quantifying a prior probability is of course inherently subjective an issue which we address later.

**How to personalize treatment effects.**

We suggest that after one identifies important potential effect modifiers (based on the criteria discussed), and quantifying the available prior knowledge, one explores if generalizability can be shown. If generalizability cannot be proven, we propose a thorough multivariable analyses to explore for which patients treatment needs to be modified.
To explore generalizability one first needs to define regions of equivalence as discussed above. After which pre-specified interaction tests can be compared against this region. In itself pre-specification does not significantly increase power to detect interactions unless proper design steps are taken (e.g., oversampling of subgroups). We suggest that, regardless of pre-specification or not, these interaction tests are deemed exploratory unless steps are made to quantify the prior evidence, ensure sufficient sample size, power, and type 1 error levels.

After determining the amount of within study heterogeneity, and assuming multiple studies exist, between study heterogeneity should be explored, for example by comparing aggregated results from different studies. However, attributing differences in treatment effects between studies to differences in baseline characteristics or study design, using for example meta-regression, may result in (ecological) bias. Therefore, significant interaction effects found in aggregated meta-analyses should always be confirmed using individual patient data (IPD) or independently replicated.

If, after performing the above analyses, absence of effect modification cannot be excluded with confidence, confirmatory analyses are needed, tailoring treatment effects towards groups or individuals. If treatment homogeneity is rejected one may be tempted to treat this as a true positive results. However, as with any discovery, replicating results is essential, hence results on interaction effects should be independently confirmed. If the results are replicated, it seems sensible to finally combined data from both the confirmatory and exploratory steps to increase precision in the subgroup specific effect
estimates of the treatment\textsuperscript{52}, and use these to tailor treatment (e.g., RR = 0.75 for diabetes patients versus RR = 0.63 in patients without diabetes).

Recently, subgroup-specific estimates based on a single variable (i.e., univariable interactions) have been criticized\textsuperscript{15,53-56}. Among other reasons, critics recognized that patients likely differ on more than one characteristic (i.e., there is unexplained treatment effect modification). A straightforward solution is to include multiple interaction tests, for example exploring whether treatment effects differ by diabetes, gender, and age. However, depending on the number of subgroups (and type, e.g., binary or not), exploring higher order interactions will increase data sparseness, which may dramatically reduce power and increase type 1 error rates\textsuperscript{10;39-42;57-62}.

To (partially) solve this, a two-step multivariable method has been suggested. First, a multivariable risk prediction model is developed, predicting the risk of the outcome if a subject is not treated\textsuperscript{63,64}. For example, using a logistic model, the predicted risk equals

\[
\logit(\hat{p}_i) = \logit(\text{Prob}[Y = 1|Z]) = \hat{\beta}_0 + \sum_{j=1}^{k} \hat{\beta}_j z_{ij} \quad \text{[equation 1]},
\]

where Z equals a \( n \times k \) matrix and Y a \( n \times 1 \) column matrix. In the second step, the predicted risk is multiplied by a relative treatment effect estimate (e.g. a risk ratio)\textsuperscript{56}. Assume, for example, that in our previous trial the multivariable 5-years predicted risk of stroke without treatment equals

\[
\frac{1}{1+e^{-\logit(\hat{p}_i)}} = \frac{1}{1+e^{2.20}} = 0.10,
\]

for a particular patient with diabetes. Based on the RR of 0.75, treating this patient will result in a predicted 5 year risk of 0.075 (i.e., 0.75 \times 0.10 = 0.075) and in an individualized RD of 0.100 – 0.075 =
0.025; in the general case the RD can be individualized using equation 2: \[ RD_i = \logit(\hat{p}_i)^{-1} * RR. \]

While this multivariable approach to subgroup analysis is indeed an improvement, one should be aware, this approach is only valid if the relative treatment effect measure (e.g., the RR) is homogenous across different levels of the predicted risk. If unknown to the researcher, the relative treatment effect measure is in fact heterogeneous, applying the above approach may falsely induce treatment effect modification on the risk difference scale. We propose that when the above approach is applied, this should be combined by the following sensitivity analysis exploring 1) if the relative effect measure is heterogeneous across the range of predicted risks and 2) to what extend the RD is truly heterogeneous across the predicted risk. Following the risk stratification approach by Kent and Hayward \(^{15}\) and others we suggest subdivide the subjects sample based on quantiles of the predicted risk (equation 1) and estimate quantile specific treatment effects (e.g., RR and RD)\(^{15}\) to explore if the treatment effects changes with increasing risk. Using this approach one can judge if the relative effect measure is fairly homogenous across the predicted risk and if individualized treatment effects on the RD scale (equation 2) agree with the quantile specific treatment effects on RD scale. As with any testing procedure one should be careful not to over interpret non-significant interaction tests results, because, as addressed before, this does not imply homogeneity. To increase power and only if quantile specific treatment estimates linearly change one could use the predicted risk as a linear term in a statistical model and include a treatment by predicted risk interaction term\(^{65}\). Depending on the amount of linearity such a model is expected to be more precise and powerful than the quantile
specific approach. A remaining issue with the above described approaches is that, typically, the predicted risk is treated as if it was observed without error, possibly erroneously decreasing the standard error of any test\textsuperscript{66}. A second more general comment is that all the discussed multivariable approaches only allow for individualized treatment effect estimates in so far as variables are related to the outcome. A strategy to include variables unrelated with the outcome in a multivariable interaction test is to use unsupervised cluster analysis to identify multivariable patient clusters, and test if treatment effectiveness differs across cluster memberships\textsuperscript{67-69}.

**Quantifying the prior probability of treatment effect modification.**
Throughout the previous sections we frequently emphasized the need to quantify the prior probability for the presence of an interaction. Here we detail what to base this prior probability on.

As stated previously, RCTs are the gold standard in intervention research. Despite this, we feel strongly against a priori deciding to quantifying the prior probability solely on RCT results. RCTs are not initiated at random. Instead, RCTs are initiated based on information from basic experiments, genetic studies, nonrandomized studies and/or previous RCTs, therefore to properly quantify the prior knowledge these sources should all be considered. Depending, however, on the potential risk of bias, taking account of the endpoints of interest, and the general potential risk of an intervention, these multiple source of prior knowledge should be reweighted. In some cases, for example, when exploring the intended effect of statins on a myocardial infarction, one may choose to weight non-RCT data by zero. This reweighing or elimination of data should obviously be
clearly presented and justified. We appreciate that this introduces a certain amount of subjectivity in analyses that may seem otherwise objective. However, this is no different than excluding RCTs at perceived high risk of bias from a meta-analysis, a thing which is customarily (although not without discussion) done in, for example, cochrane reviews.

**Summary**

In the present commentary we have argued that detecting treatment effect modification is essential to bridge the gap between results from clinical studies and treating individuals in daily practice. We addressed strategies to detect effect modification and used these in a framework to assess if there is a need for more individualized treatment effects, and estimate this in confirmatory analyses.

We conclude with the following recommendations. First, treatment effect modification should be formally assessed using interaction tests. Second, pre-specified subgroups should be selected based on biological plausibility, prevalence of the patient type and cost-effectiveness of determining the patient characteristic. Third, before tailoring treatment effects to patient subgroups one should first consider if generalizability or the absence of treatment effect modification can be proven, using e.g., an equivalence test. Fourth, for interaction tests to be anything but exploratory, these should not only be pre-specified, but include a quantification of the prior knowledge, use proper sample size calculations and sampling strategies to ensure appropriate levels of power and type 1 error rates (taking account of possible multiple testing). Finally, if after careful consideration and sufficient replication, subgroup effects are found to be consistent across different studies, this should have an impact in daily clinical practice. What is
sufficient evidence, however, should be determined on a case by case basis and depends, amongst other things, on the disease, intervention related risks and the magnitude of interaction.
Tables

Table 1 Stroke risk by exposure (X) and baseline diabetes (D) status and their interaction on different measurement scales.

<table>
<thead>
<tr>
<th></th>
<th>D = 0</th>
<th>D = 1</th>
</tr>
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<tbody>
<tr>
<td>X = 0</td>
<td>0.80</td>
<td>0.89</td>
</tr>
<tr>
<td>X = 1</td>
<td>0.50</td>
<td>0.67</td>
</tr>
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</table>

<table>
<thead>
<tr>
<th>Measure of risk difference interaction</th>
<th>Measure of risk ratio interaction</th>
<th>Measure of odds ratio interaction</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.67 − 0.50 − 0.89 + 0.80 = 0.08</td>
<td>0.67 * 0.80 = 1.20</td>
<td>0.67 * 0.80 = 1.20</td>
</tr>
<tr>
<td></td>
<td>0.50 + 0.89 = 1.39</td>
<td>0.33 * 0.20 = 0.66</td>
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<tr>
<td></td>
<td></td>
<td>0.89 + 0.50 = 1.39</td>
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<tr>
<td></td>
<td></td>
<td>0.11 * 0.50 = 0.05</td>
</tr>
</tbody>
</table>
Figure captions

Figure 1. Examples of equivalence testing using confidence intervals*.
*Based on Jones et al. 33.

Figure 2. Empirical power of two test for equivalence of treatment effect modification*.

* The dashed line with a square symbol indicates power for an equivalence test using an interaction effect, the dashed dotted line with the circle symbol indicates power for the equivalence test based on subgroup specific effects. Simulated results were based on a scenario (1,000 replications) with subjects treated or untreated $j = \{1,0\}$ and exposed or unexposed to a potential effect modifier $i = \{1,0\}$, with the endpoint incidence equaling $r_{ij} = \{0.20, 0.15, 0.15, 0.10\}$, and each group of $ij$ subjects occurring 1,000 times.
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