HER2-enriched subtype and novel molecular subgroups drive aromatase inhibitor resistance and an increased risk of relapse in early ER+/HER2+ breast cancer

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Summary

Background Oestrogen receptor positive/ human epidermal growth factor receptor positive (ER+/HER2+) breast cancers (BCs) are less responsive to endocrine therapy than ER+/HER2- tumours. Mechanisms underpinning the differential behaviour of ER+HER2+ tumours are poorly characterised. Our aim was to identify biomarkers of response to 2 weeks’ presurgical AI treatment in ER+/HER2+ BCs.

Methods All available ER+/HER2+ BC baseline tumours (n=342) in the POETIC trial were gene expression profiled using BC360™ (NanoString) covering intrinsic subtypes and 46 key biological signatures. Early response to AI was assessed by changes in Ki67 expression and residual Ki67 at 2 weeks (Ki67 2wk). Time-To-Recurrence (TTR) was estimated using Kaplan-Meier methods and Cox models adjusted for standard clinicopathological variables. New molecular subgroups (MS) were identified using consensus clustering.

Findings HER2-enriched (HER2-E) subtype BCs (44.7% of the total) showed poorer Ki67 response and higher Ki67 2wk (p<0.0001) than non-HER2-E BCs. High expression of ERBB2 expression, homologous recombination deficiency (HRD) and TP53 mutational score were associated with poor response and immune-related signatures with High Ki67 2wk. Five new MS that were associated with differential response to AI were identified. HER2-E had significantly poorer TTR compared to Luminal BCs (HR 2.55, 95% CI 1.14–5.69; p=0.0222). The new MS were independent predictors of TTR, adding significant value beyond intrinsic subtypes.

Interpretation Our results show HER2-E as a standardised biomarker associated with poor response to AI and worse outcome in ER+/HER2+. HRD, TP53 mutational score and immune-tumour tolerance are predictive biomarkers for poor response to AI. Lastly, novel MS identify additional non-HER2-E tumours not responding to AI with an increased risk of relapse.

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Introduction

Human epidermal growth factor receptor 2 positive (HER2+) breast cancer (BC) has been associated with an aggressive phenotype and poor patient outcome.1 However, the introduction of HER2-targeted therapies dramatically changed the prognosis of these patients.
Research in context

Evidence before this study

AI treatment is the standard of care and most effective therapy for post-menopausal women with early oestrogen receptor positive (ER+) breast cancer (BC). ER+ BC tumours that also over-express HER2 are very heterogeneous with several treatment options but variable responses to the different available drugs. Prior studies have shown that ER+/HER2+ BC show limited antiproliferative response to endocrine therapy and thus, are at a higher risk of recurrence. This may be PgR dependent, as many of those tumours do express low PgR levels. In addition, they have lower response rates to anti-HER2 targeted therapy compared to ER-/HER2+ tumours. Most studies investigating mechanisms of resistance to endocrine therapy have been performed in ER+/HER2- disease and are not well understood in HER2+ BC. Although endocrine-related gene expression has been previously associated with good response to aromatase inhibitors and high levels of ERBB2 with poor response, there is an overall lack of optimal biomarkers to pair with the optimal treatment for each patient within ER+/HER2+ BC. As such, identifying robust molecular features and defining novel subgroups based on tumour biology is essential to identify the most adequate treatment strategies for this particular BC subgroup.

Added value of this study

This study establishes HER2-enriched intrinsic subtype as one of the main components driving poor response to AI and higher risk of relapse in ER+/HER2+ BC. Our results indicate the importance of molecular subtyping of BC beyond the standard HR and HER2 assays. Beyond the intrinsic subtypes, ERBB2, DNA damage repair signaling, p53 mutant surrogate signature and immune-tumour tolerance related signatures are also associated with resistance to treatment.

We also identified five new single gene based molecular subgroups that can distinguish HER2-E and Luminal tumours responding or not to AI treatment and at a higher risk of relapse. Molecular subgroups characterised with high expression of immune related features drive an intrinsic lower risk of relapse despite predicting poor response to AI, while higher levels of ERBB2 and extracellular matrix related genes lead to worse outcome.

Implications of all the available evidence

Firstly, the worse response to treatment and poorer outcome of HER2-enriched BC tumours highlights the importance of development of resistance to AI but while anti-HER2 therapy is generally given for no more than 1 year to primary BC patients, ET is given for at least 5 years. Thus, any residual HER2+ disease after the end of the HER2-targeted therapy remains at risk of an incomplete endocrine response.

The PeriOperative Endocrine-Therapy for Individualised Care (POETIC) trial is the framework used to study endocrine resistance mechanisms in a large set of ER+/HER2+ BC patients. In the context of POETIC, we hypothesised that resistance mechanisms to ET are driven by baseline genomic features. Gene expression profiles at baseline were assessed and the key genomic characteristics were tested for association with response and the natural history of the disease. Despite the improvement, long-term follow-up data indicate that approximately 15–23% of patients in early stage, still develop recurrent disease.

Fifteen percent of all BC overexpresses HER2 and approximately 50% of these are also classified as hormone receptor positive (HR+), which confers substantial differences in biology and clinical outcome from HR+/HER2- disease. HR+/HER2+ BCs are molecularly heterogeneous and around 30% of them are HER2-Enriched (HER2-E). This subtype is characterised by a high HER2/EGFR pathway activation, increased proliferation and an immune-activated stroma with elevated tumour infiltrating lymphocytes. It has a lower expression of luminal-related genes, than the Luminal A and B subtypes, potentially benefiting greatly from anti-HER2 therapies but poorly from endocrine therapy (ET).

Resistance to endocrine therapies has been mainly studied in HR+/HER2- BC and includes down-regulation of oestrogen receptor (ER) expression, altered expression of ER co-regulators, presence of ER mutations, ligand-independent activation of ER and co-activators by growth factor receptor kinases. However, those mechanisms might differ between HER2+ and HER2- tumours, in part due to the differential distribution of intrinsic subtypes within each BC subgroup. HER2-targeted therapies might be felt to negate the importance of development of resistance to AI but while anti-HER2 therapy is generally given for no more than 1 year to primary BC patients, ET is given for at least 5 years. Thus, any residual HER2+ disease after the end of the HER2-targeted therapy remains at risk of an incomplete endocrine response.
to AI, as measured by residual levels and changes of Ki67 after two weeks of treatment, and with clinical outcome. We sought to develop predictive signatures of AI response and address the clinical challenge of identifying patients who are likely to benefit from each of the specific therapies.

Methods

Patients and samples
All available ER+/HER2+ BC tumours from the POETIC trial in which patients were assigned to 2 weeks of peri-surgical AI or no AI (control) were included in this study. A consort diagram of the study is shown in Supplementary Figure S1. Ki67 staining of 2-week samples from the control group was restricted to a randomly selected subset due to the minimal expected change on Ki67 from baseline to surgery. In summary, of 470 ER+/HER2+ patients included in POETIC, we obtained successful results for 342 patients.

RNA extraction
RNA was extracted from three adjacent macro-dissected 10µm formalin-fixed paraffin-embedded (FFPE) sections from the baseline block of the patients included in the study. The ROCHE High Pure miRNA isolation kit (Roche, Basel, Switzerland) was used following SOP M027 from The Cancer Genome Atlas (TCGA) Program developed by the Biospecimen Core Resource (BCR) at Nationwide Children’s Hospital in Columbus, Ohio. Quantification was done using high sensitivity RNA Qubit assays (Thermo Fisher Scientific, Carlsbad, CA).

Gene expression profiling
Gene expression of 758 genes was assessed using the NanoString nCounter Platform (Nanostring Technologies, Seattle, WA) Breast Cancer 360™ codset (BC360) covering intrinsic subtypes and 46 key biological signatures (Supplementary Table S1). 150ng of RNA was run and processed on a NanoString nCounter™ FLEX Analysis System according to manufacturer’s instructions. NanoString raw data was normalised by NanoString according to the BC360 pipeline using 18 house-keeping genes.

Immunohistochemistry
ER status was measured locally and was centrally reviewed by immunohistochemistry (IHC). HER2 status was measured locally using IHC and/or fluorescence in situ hybridisation (FISH). Ki67 proliferation rate was obtained by IHC in FFPE tissues using the MIB-1 antibody (M7240, DAKO UK, RRID: AB_2631211). Ki67 has been validated in our laboratory previously and we are part of the International Ki67 in Breast Cancer Working Group.

Outcomes
The primary endpoints of this study were based on Ki67 as a measure of tumour’s resistance to AI. Two Ki67 endpoints were used: 1) Ki67 change was calculated as the difference between Ki67 expression at surgery and baseline (relative change) and was categorised into Ki67 response categories defined as percentage-changes from baseline to surgery: poor response (PR) (reduction <50%), intermediate response (IR) (50–75%) and good response (GR) (>75%). This reflects the antiproliferative response to AI treatment which relates to the treatment benefit. 2) Residual Ki67 at 2-week timepoint (Ki672wk) High (≥10%) and Low (<10%) which correlated to the residual risk after AI treatment. The secondary endpoint was time to recurrence (TTR) (local and metastatic) to evaluate the prognostic significance of the molecular characteristics analysed.

Statistical analysis
Statistical analysis was performed using the R software (version 3.6.3). P values were considered significant if lower than 0.05. Wilcoxon tests were applied in unpaired comparisons and Kruskall-Wallis tests in all multiple comparisons in both treated and control tumours. Spearman Rank correlation was used to explore the correlation between genes or signatures. Logistic and ordinal regression models were performed to identify signatures significantly associated with 2-week Ki67 and Ki67 response categories respectively. Multiple testing correction was undertaken by the Benjamini & Hochberg (FDR) method. Significance analysis of microarrays (SAM analysis) was performed in multiclass setting to compare the Ki67 response categories and in the unpaired Two Class to compare extreme response classes (GR and PR) and Ki672wk High versus Low. Hierarchical clustering of the gene expression profiles that were identified by SAM analysis was also performed. Consensus clustering was used to identify new molecular subgroups and their association with Ki67 response categories and outcome was tested. Controls and treated patients were included to obtain the subgroups, but only the treated patients with Ki67 available, were used to assess their predictive value for AI resistance.

TTR was measured as time from randomisation to local, regional, or distant tumour recurrence or death from breast cancer without previous notification of relapse. Second primary cancers and intercurrent deaths were censored. TTR was estimated using Kaplan-Meier methods and Cox models. Multivariable Cox-regression models were adjusted for standard post-surgery clinicopathological variables: grade, tumour size, nodal status and age. We included age as the main driver/surrogate for the adjuvant treatment choice as most patients ≥70 years old did not receive...
chemotherapy or trastuzumab (67.2%, 82/122) compared to patients <70 years (14.5%, 31/211). The independent prognostic value of those gene-expression based variables with differential survival in the univariate analysis were assessed. Both controls and treated patients were included in the survival analysis. The assumptions evaluated were: 1. that the dependent variable was ordered; 2. that one or more of the independent variables were continuous, categorical or ordinal; 3. that no multi-collinearity i.e. independent variables were independent from each other and the proportional odds. And 4. that the tests were done for the proportional odds assumptions for the ordinal logistic regression analysis for single gene and signature with the Ki67 change categories GR/IR/PR. All the assumptions were met without any assumptions’ violations. Patients with missing data were excluded. Only one patient had a missing variable (which constituted only 0.29% of the entire cohort).

**Ethics**

The POETIC trial was approved by the London–South East Research Ethics Committee (reference 08/H1102/37) and adopted by the Declaration of Helsinki. Patients provided written informed consent to molecular analysis of their samples for research purposes.

**Role of funding source**

Funders did not have any role in study design, data collection, data analysis, interpretation or writing of report.

**Results**

**Patient clinicopathological characteristics**

In this study, 342 ER+/HER2+ patients with baseline gene expression were included: 237 AI-treated and 105 untreated controls (Supplementary Figure S1). The demographics were well balanced between both groups (Supplementary Table S2). In summary, 93.3% of the tumours were ductal, 48.2% were grade 2 and 38.3% grade 3. At surgery 54.7% had a tumour diameter between 2 and 5 cm and 47.4% had positive nodal status. 60.5 percent of patients received adjuvant chemotherapy and trastuzumab and 98.2% of patients were treated with adjuvant ET.

**PAM50 subtypes and Ki67 endpoints**

We evaluated whether intrinsic subtypes could predict response to ET. In the entire subset of patients, 44.7% of tumours classified as HER2-E, 36.3% Luminal B, 17.2% Luminal A and 18% Basal-Like. In addition, the proportion of intrinsic subtypes was comparable between control and treated groups with the controls slightly enriched with HER2-E (54% vs 41.4%) and reduced Luminal A tumours (8-0% vs 21.1%), not statistically significantly different (p>0.05). Within the treated subgroup, 31% achieved GR, 22-5% were IRs and 46.5% PRs, while 52-0% had Ki672wk High and 48.0% Low, respectively.

As expected, most control tumours were classified as PRs (96-0%) and Ki672wk High (96-0%) (Table 1). In the treated group, there was a significant change in Ki67 values in all subtypes (Figure 1A), except for Basal-like, possibly due to the small sample number. Overall, the HER2-E subtype was associated with poorer response to AI compared to non-HER2-E, evaluated as Ki67 response category and residual Ki67. HER2-E tumours were associated with higher grade (68.6% grade 3 in HER2-E vs 30.9% in luminals, p<0.0001, Fisher test) and larger tumours (50-3% >3 cm in HER2-E vs 39.1% in luminals, p<0.0001, Fisher test). However, the association of the HER2-E subtype and Ki67 response remained significant in each of the categories.

**Table 1:** Distribution of the 4 intrinsic subtypes within patients with Ki67 data and by Ki67 categories

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<tr>
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<th>TREATED</th>
<th>CONTROLS</th>
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<tr>
<td></td>
<td>Arm</td>
<td>HER2-E</td>
</tr>
<tr>
<td></td>
<td>All (100.0%)</td>
<td>95 (42.1%)</td>
</tr>
<tr>
<td></td>
<td>Basal (3.1%)</td>
<td>3 (1.3%)</td>
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<tr>
<td></td>
<td>HER2-E</td>
<td>18 (6.2%)</td>
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<tr>
<td></td>
<td>LumA</td>
<td>63 (22.9%)</td>
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<tr>
<td></td>
<td>LumB</td>
<td>56 (19.9%)</td>
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Chi-squared 27.69, P<0.0001

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<th>Ki67 Response Categories</th>
<th>Chi-squared 27.69, P&lt;0.0001</th>
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<tbody>
<tr>
<td></td>
<td>GR</td>
<td>0 (0.0%)</td>
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<tr>
<td></td>
<td>IR</td>
<td>2 (4.0%)</td>
</tr>
<tr>
<td></td>
<td>PR</td>
<td>9 (18.0%)</td>
</tr>
</tbody>
</table>

Fisher’s exact test P=0.05

**Abbreviations:** HER2-E, HER2-Enriched Subtype; LumA, Luminal A subtype; LumB, Luminal B subtype; GR, Good response; IR, Intermediate response, PR, Poor response; Ki672wk, Ki67 at 2 weeks timepoint.
within those variables. In the control patients with Ki67 data, no significant changes of Ki67 were observed amongst subtypes (Figure 1B). These findings suggest that HER2-E might be one of the main components driving poor early response to AI in ER+/HER2+ BC tumours.

**Signature expression and Ki67 endpoints**

We then evaluated the association of other biological molecular features (46 signatures) with the defined Ki67 endpoints (Supplementary Table S3). High expression of endocrine related signatures such as ESR1, ER-Signaling, FOXA1 and PgR as well as Luminal A and B correlation coefficient scores were associated with GR and Low Ki672wk (OR 0.55–0.82; FDR<0.0001, ordinal logistic regression model), while high ERBB2, Basal-like and HER2-E correlation coefficient scores were associated with PR and High Ki672wk (OR 1.52–12.31; FDR<0.0001, ordinal logistic regression) (Figure 2). Noteworthy, the high expression of apoptosis signature (pro-apoptotic) was associated with GR (OR 0.26; 95% CI 0.12–0.54; FDR<0.0017, ordinal logistic regression) whilst high DNA-damage repair signatures such as the homologous recombination deficiency (HRD), hypoxia, and the TP53 mutational status’ surrogate signature were associated with PR (OR 2.18–2.65, FDR<0.0001–<0.0059, ordinal logistic regression). Additional high expression of signatures involved in immune-checkpoint component and tumour immunity such as IDO1, IFN Gamma, PD-L1 and Tumour Inflammation Signature (TIS) as well as the genomic risk score were associated with High.

![Figure 1. Ki67 changes from baseline to surgery stratified by intrinsic subtype at baseline within a. Treated and b. Controls. Abbreviations: HER2-E, HER2-Enriched Subtype; LumB, Luminal B subtype; LumA, Luminal A subtype.](image-url)
Figure 2. Signatures significantly associated with differential response to AI. a. Ordinal regression models in treated patients with Ki67 data (n=227) coloured by FDR for Ki67 response categories and b. Logistic regression models for Ki672wk. Abbreviations: AI, Aromatase inhibitors; logOR, log Odds Ratio; FDR, False discovery rate; HER2-E, HER2-Enriched Subtype; LumB, Luminal B subtype; LumA, Luminal A subtype.
A coefficient scores were associated with Low Ki672wk (OR 1.67–1.49, FDR < 0.0001–0.0084, logistic regression). Following prior evidence, we also assessed the correlation of the different signatures with Ki67 > or < than 20% to validate our findings. The same signatures were associated with differential response to AI based on this Ki67 threshold (Supplementary Table S4).

To assess the above associations according to the different subtypes, we tested them in the HER2-E and Luminal subtypes separately. The signatures were not significantly differently expressed between response categories amongst HER2-E tumours for any of the two Ki67 endpoints (Supplementary Figure S2A). However, in the Luminal tumours, proliferation, **TP53** mutational score, genomic risk and the Basal-like and HER2-E coefficient scores were associated with **High** Ki672wk (OR 1.15–16.84; FDR = 0.0005–0.01, logistic regression), while high expression of AR signature and the **Luminal A** coefficient scores were associated with **Low** Ki672wk (OR 0.25–0.77; FDR = 0.0019–0.013, logistic regression) (Supplementary Figure S2B). Thus, beyond HER2-E intrinsic subtype: **ERBB2**, **TRP53** mutational status, HRD and some immune-related signatures were also associated with early resistance to AI.

**Single gene expression and Ki67 endpoints**

Multiclass SAM analysis of single gene expression for the three Ki67 response categories (GR/IR/PR) (Δ = 0–26, FDR < 0.05) identified 8 genes with significantly different expression amongst groups (p < 0.05). High expression of **GRB7** and **ERBB2** were associated with PR while high expression of others such as **IGF1R**, **ESR1**, **CHAD** and **BCL2**, were associated with **GR** (Figure 3A). Two unpaired class SAM analysis for **GR** versus PR (Δ = 1, FDR < 0.05) showed the association of 31 genes with **GR** (p < 0.05, Wilcoxon test) (Figure 3B). Two class unpaired SAM analysis with **Ki672wk** categories: **High** versus **Low** (Δ = 1–63, median FDR = 0) identified 128 genes associated to **Low** Ki672wk, including genes involved in **PI3K/AKT**, **MAPK** and oestrogen signaling and 83 genes associated to **High** Ki672wk, including genes involved in immune-checkpoint component, proliferation and cell-cycle regulation (Figure 3C).

To further investigate the biological differences between early AI responders and resistant tumours we evaluated the **ESR1** gene expression levels amongst the four intrinsic subtypes. The highest levels of **ESR1** were seen in Luminal B tumours. Noteworthy, within the Luminal A tumours a group of patients showed lower levels of **ESR1** were associated with higher levels of **ERBB2** signaling, especially when compared with other subtypes (Supplementary Figure S3).

Overall, at a single gene expression level, ER signaling-related genes drive responses to AI treatment while HER2 and immune-related genes associates with early resistance.

**Identification of new molecular subgroups predicting Ki67 endpoints**

Using consensus clustering, we identified 5 novel molecular subgroups of samples based on single-gene expression that are associated with different Ki67 response and Ki672wk (Figure 4 and Supplementary Table S5). Interestingly, these new molecular subgroups divided mainly HER2-E samples with lower response to AI, into three groups with differential expression of molecular features such as **ERBB2**, ER signaling or immune-related pathways.

Figure 4 shows the molecular features of the clusters (C): C1, C2 and C3 were characterised by PR to AI (60/111, 54% and **high** Ki672wk **82/111, 73%**) and an enrichment of HER2-E subtype (79/111; 71% and **Luminal B** tumours (21/111; 18%)) (Supplementary Figure S4A). C1 (29% of the total) showed higher levels of immune and chemokine-related genes and lower levels of **ESR1** (Supplementary Figure S4B). C2 (4.2% of the total) had higher levels of extracellular matrix organization (ECM) related genes, lower levels of **ESR1** and the highest levels of **ERBB2** (Supplementary Figure S4C), also observed when distributed by subtypes. C3 (5.7% of the total) had low expression of most genes but a slight upregulation of genes covering DNA-damage repair deficiency. C4 (22.1% and C5 (28.8%) were characterised by GR to AI and an enrichment of luminal tumours (Luminal B 81/165; 49% and Luminal A 57/165; 34%) (Supplementary Figure S4A and Supplementary Table S5). Both luminal subgroups overexpressed ER signaling-related genes. However, while C4 had the highest levels of **ESR1** (Figure 5C), C5 was enriched with genes involved in **MAPK/PI3K** and **RAS** signaling. Figure 4 also shows the signatures’ expression within each of the single gene-based molecular subgroups, thus confirming the biologic distinctiveness of each of the new molecular subgroups.

**Time to recurrence analysis**

Median follow-up was 62.9 months (IQR 58.1–74.1). Univariate analysis was performed to evaluate the prognostic value of each individual genomic feature analysed. There was no overall significant difference of TTR between intrinsic subtypes when all the subgroups were compared together (Figure 5A). However, HER2-E had significantly poorer TTR (HR 2.14; 95% CI 1.11–4.17; p = 0.022, univariate regression model) compared to luminal tumours (Figure 5B). C2, which is characterised by low expression of immune-related genes and the highest **ERBB2** expression, showed a significantly higher risk of recurrence compared to the rest of the molecular subgroups (HR 3.05; 95% CI 1.31–7.15; p = 0.01, univariate regression model) (Figure 5C).

We performed a series of cox regression models for multivariable survival analysis including standard clinicopathologic factors along with different molecular subgroups for TTR and compared the changes of chi-
Figure 3. Hierarchical clustering using significant genes from SAM analysis of single gene expression for the different Ki67 endpoints in treated patients. a. Supervised hierarchical clustering of the multiclass SAM analysis for the three Ki67 response categories (GR/IR/PR) (n=226). b. Supervised hierarchical clustering of the two unpaired class SAM analysis (n=226). c. Supervised hierarchical clustering of the genes selected by two class unpaired SAM analysis for Ki672wk Categories: High vs Low (n=227): 211 significant genes, 128 associate with low and 83 with high residual Ki67. Abbreviations: HER2-E, HER2-Enriched Subtype; LumB, Luminal B subtype; LumA, Luminal A subtype; Ki672wk, Ki67 at 2 weeks timepoint.
Figure 4. Five new molecular subgroups were identified based on consensus clustering of tumours using expression of 758 genes. Top panel illustrates single gene expression of all genes in treated patients with Ki67 data only (n=226), separated by the 5 novel molecular subgroups. Clusters of genes are annotated according to their functionality. Bottom panel illustrates the expression levels...
squared values between them to assess the added value of the different models (Supplementary Table S6). HER2-E subtype remained as an independent predictor of higher risk of relapse (HR 2.55, 95% CI 1.14–5.69; $p=0.022$, multivariable regression model). C2 was also an independent predictor of shorter TTR compared to C1, indicating that this model adds significant value beyond intrinsic subtypes (Likelihood ratio test, $p=0.0025$). (Supplementary Figure S5).

To explore further the role of adjuvant HER2-targeted treatment and chemotherapy on survival, we replicated the same analysis but separately for those patients who had received both of these treatments and those that did not (Supplementary Figure S6). Noteworthy, among patients who had received adjuvant trastuzumab plus chemotherapy, there were no significant differences in TTR observed between HER2-E and other subtypes (Supplementary Figure S6a). However, among patients who had not received trastuzumab and chemotherapy, HER2-E was still significantly associated with worse outcome compared to non-HER2-E (Supplementary Figure S6a). For the new molecular subgroups, the survival outcome within the new molecular subgroup 2 was worse compared to the others in those patients receiving adjuvant anti-HER2 treatment. In addition, in the multivariable models, the new molecular subgroups remained as independent predictors of TTR restricting to patients who received trastuzumab plus chemotherapy (HR 3.95; 95% CI 1.04–14.94; $p=0.0043$, multivariable regression model).

Finally, a series of multivariable analyses for TTR of signature expression adjusted for the clinicopathological factors showed that several immune related signatures (HR 0.50–0.78, nominal $p<0.0001–0.0430$, multivariable regression model) as well as the apoptosis signature (HR 0.36, 95% CI 0.15–0.38; $p=0.024$, multivariable regression model) were independent predictors of better TTR. By contrast, claudin-low signature showed independent worse prognostic value (HR 1.26, 95% CI 1.06–1.56; $p=0.031$, multivariable regression model) (Supplementary Table S7). A final model including all identified features was not performed due to the high collinearity amongst the main significant signatures (Supplementary Figure S7).

**Discussion**

AI treatment is the standard of care and most effective therapy for post-menopausal women with early ER+ BC. However, ER+ tumours that also over-express HER2, show limited response to ET and thus, are at a higher risk of recurrence. Most studies performed in ER+/HER2+ BC have focused on resistance to anti-HER2 targeted therapy while mechanisms of resistance to ET are not well understood within this subgroup. This study investigated the predictive and prognostic value of molecular features at baseline in ER+/HER2+ BC treated with perioperative AI. Prior studies have revealed that HER2-E subtype is a predictor of higher sensitivity to anti-HR2 targeted therapy but worse outcome than other subtypes such as luminal tumours. Nevertheless, the role of the intrinsic subtypes in response to ET has not been well established yet. A prior single-arm, multicentric study (PerELISA) included 65 postmenopausal patients with HR+/HER2+ operable patients receiving 2 weeks of letrozole and then undergoing to re-biopsy for Ki67 evaluation. This study reported the association of PAM50 intrinsic subtyping with molecular responders (Ki67 relative reduction $>20\%$ from baseline). 92% of responders were luminal A and B versus 44% HER2-E and basal-like ($P<0.001$). These results were aligned with ours; however, this is a small, single arm, non-randomised study, while POETIC was a randomised trial, enabling us to evaluate the predictive value of the intrinsic subtyping. Within PerELISA trial patients classified as molecular responders continued letrozole and started trastuzumab-pertuzumab for five cycles. They also reported that the pCR rate was significantly higher in HER2-E than in other subtypes ($45.5\%$ versus $13.8\%$, $P=0.042$). There was no association test of survival outcome reported, nevertheless, these results are of clinical significance as this provide a potential strategy for de-escalating the treatment management.

Our results identified HER2-E subtype as a predictive biomarker of resistance to AI in ER+/HER2+ BC with an additional higher risk of relapse. These findings highlight two aspects. First, they suggest the potential need of treatment intensification for HER2-E ER+/HER2+ BC tumours with intensive anti-HER2 targeted therapy. Second, the higher sensitivity to AI and good prognosis associated with luminal tumours, in particular with Luminal A, provides a rationale for testing de-escalation approaches, such as the reduction of anti-HER2 blockade duration or avoiding chemotherapy, previously suggested. A related finding was published in 2021 as part of the ADAPT HER2+ trial in which the pooled TDM1 arm of patients with HER2-E subtype had a higher pCR, which was not observed in the trastuzumab arm.

**Abbreviations:** HER2-E, HER2-Enriched Subtype; LumB, Luminal B subtype; LumA, Luminal A subtype; GR, Good responders; IR, Intermediate responders, PR, Poor Responders; Ki672wk, Ki67 at 2 weeks timepoint, ER, Oestrogen receptor; HRD, Homologous recombination deficiency ECM, extracellular matrix; TIS, Tumour Inflammation Signature; TIGIT, T cell immunoreceptor with Ig and ITIM domain; MHC, Major Histocompatibility complex APM, Antigen Processing Machinery, IFN-gamma, interferon gamma.
a. Kaplan Meier curves according to all intrinsic subtype at baseline.

b. Kaplan Meier curves according to intrinsic subtype at baseline: HER2-E vs Luminals (A and B), excluding Basal-like tumours

c. Kaplan Meier curves according to the 5 new molecular subgroups

**Figure 5.** Kaplan Meier curves for TTR according to a. all PAM50 intrinsic subtype at baseline; b. HER2-E vs luminals and c. the new molecular subgroups. Abbreviations: TTR: Time to recurrence; HER2-E, HER2-enriched; LumB, Luminal B.
Nowadays, the biggest question is how to select patients for escalation and de-escalation strategies. Recent “chemotherapy-free” neoadjuvant studies have shown that the combination of HER2-E subtype with high ERBB2 mRNA expression may identify patients with HER2+ early BC with higher sensitivity to double HER2-blockade. However, only one third of ER+/HER2+ BC patients are HER2-E/ERBB2 high and their role as prognostic biomarkers is still unknown. Using consensus clustering, we identified new molecular subgroups based on single gene expression at a higher risk of relapse beyond HER2-E subtype. Results from the stratified analysis by adjuvant treatment (chemotherapy plus anti-HER2 treatment yes vs no) showed similar TTR despite intrinsic subtype in those patients receiving trastuzumab plus chemotherapy. This highlights the hypothesised notion that patients with HER2-E subtype benefit the most from anti-HER2 treatment. The independent prognostic value for the new molecular subgroups for TTR in patients who received trastuzumab plus chemotherapy suggests the potential clinical utility of these new molecular subgroups to identify a set of ER+/HER2+ patients, beyond intrinsic subtypes, who may benefit from emerging treatment such as CDK4/6 inhibitors.

Noteworthy, tumours characterised by an enrichment of immune features showed the lowest risk of recurrence despite immune features being predictive markers of resistance to AI and poor outcome in ER+/HER2- BC. Due to its intrinsic good prognosis, BC patients within this molecular subgroup showing an enrichment of immune characteristics could potentially benefit the most from a de-escalating approach, using for example 6 month of trastuzumab plus AI without chemotherapy followed by 5 years of AI treatment alone. Several trials in HER2+ BC have shown that tumours with higher baseline tumour infiltrating lymphocytes (TILs) and other immune features achieve higher pathological complete response rates and improved event-free survival. Our study confirms the association of higher expression of immune-features with a very low risk of recurrence in ER+/HER2+ BC. By contrast, tumours with higher levels of ERBB2 and lower associated immunity had a significantly higher risk of relapse, indicating that these patients may benefit from an intensified anti-HER2 treatment, using for example double anti-HER2 blockade or adjuvant TDM1. Therefore, these new molecular subgroups might be essential to identify candidates for new escalating and de-escalating strategies.

The role of tumour immunity in response to ET has been mainly studied in ER+/HER2- BC disease, with higher expression of genes involved in immune enrichment and targetable immune checkpoint components being correlated with higher risk such as Luminal B tumours. Our study shows that higher tumour immunity might also be a key driver of early resistance to ET in ER+/HER2+ BC. However, in this subgroup the association of higher expression of immune-features and lower risk of recurrence indicates the different role of tumour immunity between HER2+ and HER2- disease and may suggest a de-escalation approach.

Furthermore, our study shows other striking molecular associations such as high expression of HRD and TP53 signature predicting poor response to AI, and higher apoptosis signaling being associated with good response and survival. The association of DNA damage repair defects with resistance to ET has been previously reported. In addition, the inhibition of poly (ADP-ribose) polymerase-1 (PARP), has shown anti-tumour effects with a strong synergism and good tolerance in combination with anti-HER2 targeted therapy or ET in vitro and in phase I/II trials independently of DNA repair deficiency. TP53 mutational status has also been previously linked with poor survival and overexpression of HER2, although its role in response to AI is still unclear. Previous data has shown that some molecular features related to apoptosis can be predictive of adjuvant benefit from ET. Based on our results, the association of high expression of some of the signatures reported above such as immune-related signatures and TP53 mutational score with poor AI response could be explained by the high correlation with the HER2-E subtype.

Our study has two main limitations. Firstly, we only analysed gene expression for 758 genes and intrinsic subtypes profiles using the BC360 platform and other important pathways may have been missed. Secondly, clinical practice is currently different to that implemented in the recruited POETIC patients as high-risk tumours would be receiving additional pertuzumab and further anti-HER2 agents such as TDM1. It is noteworthy that one third of the patients in our cohort did not receive any adjuvant treatment apart from ET due to their advanced age, but we adjusted the survival analysis for age as a main surrogate of treatment choice. The major strengths of our study are that to our knowledge this is the largest cohort investigating response to pre-surgical AI in ER+/HER2+ subgroup in a real-world cohort which has a unique value to assess global gene expression data at baseline as defined in the clinical practice. Lastly, we were able to analyse both the molecular features associated to mechanisms of resistance and their prognostic value.

In conclusion, HER2-E subtype and ERBB2 play a crucial role in ER+/HER2+ BC, driving resistance to ET and a higher risk of recurrence. Beyond them, new molecular subgroups enable the identification of patients at a higher risk of relapse. Altogether, the combination of these biomarkers would lead to a better tailoring of treatment strategies, including escalation and de-escalation approaches, to improve resistance to treatment in early BC. The appropriate new strategies need to be addressed in prospective clinical trials.
Contributors
MB and ELK designed, conducted part of the research work, analysed and verified the data and wrote the original draft of the manuscript. GM conducted part of the research work. HT did data analysis and interpretation. LK participated in study design, data collection and data interpretation. ES supervised some of the analysis of the data. AA and MH conducted part of the research work. HX participated in data analysis for manuscript revision. CH recruited patients to the POETIC trial and provided tissue. AS participated in POETIC study design, POETIC trial management group and clinical recruitment. JFR participated in POETIC trial design and reviewing the manuscript. IS participated in the design of original POETIC trial and was chief investigator of it. JMB has provided oversight and guidance for trial management, statistics and data interpretation throughout the trial including insights into the translational research project presented here. She reviewed and contributed to the manuscript. MD was involved in the investigation and visualisation of the study. MCUC designed and led the study. She also led the conceptualisation, methodology, resources and funding acquisition, data curation and statistical analysis plan, supervision. All authors made substantial contributions to the manuscript, revised the manuscript critically, gave their final approval and had full access to all the data and accept responsibility to submit for publication.

Data sharing statement
De-identified data will be made available to other researchers on request, subject to approval of a formal data access request in accordance with the ICR-CTSU data and sample access policy. Gene expression data is available on request by contacting poetic-icrcitsu@icr.ac.uk. The ICR-CTSU supports the wider dissemination of information from the research it does, and increased cooperation between investigators. Trial data is collected, managed, stored, shared, and archived according to ICR-CTSU Standard Operating Procedures in order to ensure the enduring quality, integrity, and utility of the data. Formal requests for data sharing are considered in line with the Institute of Cancer Research Clinical Trials and Statistics Unit (ICR-CTSU) procedures with due regard given to funder and sponsor guidelines. Requests are via a standard proforma describing the nature of the proposed research and extent of data requirements. Data recipients are required to enter a formal data sharing agreement which describes the conditions for release and requirements for data transfer, storage, archiving, publication and intellectual property. Requests are reviewed by the Trial Management Group (TMG) in terms of scientific merit and ethical considerations including patient consent. Data sharing is allowed if proposed projects have a sound scientific or patient benefit rationale as agreed by the TMG and approved by the Trial Steering Committee as required. Restrictions relating to patient confidentiality and consent will be limited by aggregating and anonymising identifiable patient data.

Declaration of interests
MB reports grants from Fundacion Alonso Martin Escudero and has a patent filed for the novel molecular subgroups identified in this paper. ELK has a patent filed for the novel molecular subgroups identified in this paper. HT and LK report grants from Cancer Research UK, during the conduct of the study. JMB reports grants from Cancer Research UK, during the conduct of the study; grants and non-financial support from AstraZeneca, Merck Sharp & Dohme, Puma Biotechnology, Clovis Oncology, Pfizer, Janssen-Cilag, Novartis, Eli Lilly, and Roche, outside the submitted work. MD receives consulting fees from AstraZeneca, Lilly, Roche, Radius, H3 Biomedicine and G1. He also receives honoraria from Nanostring. M.C.U.C. has a patent for Breast Cancer Classifier: US Patent No. 9,631,339 with royalties paid and receive research funding from NanoString Technologies and Veracyte advisory role. And has a patent filed for the novel molecular subgroups identified in this paper. All other authors declare no competing interests. The information in this manuscript is subject to a pending patent application.

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Supplementary materials
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