Whole-genome analysis informs breast cancer response to aromatase inhibition

Matthew J. Ellis1,2,3*, Li Ding4,5*, Dong Shen4,5*, Jingqin Luo3,6, Vera J. Suman7, John W. Wallis4,5, Brian A. Van Tine8, Jeremy Hoog9, Reece J. Goiffon8,9,10, Theodore C. Goldstein11, Sam Ng11, Li Lin1, Robert Crowder1, Jacqueline Snider1, Karla Ballman2, Jason Weber1,8,12, Ken Chen13, Daniel C. Koboldt4,5, Cyriac Kandoth4,5, William S. Schierding4,5, Joshua F. McMichael4,5, Christopher A. Miller4,5, Charles Lu4,5, Christopher C. Harris4,5, Michael D. McLellan4,5, Michael C. Wendl4,5, Katherine DeSnyder4,5, D. Craig Allred3,14, Laura Esserman13,15, Gary Unzeitig16, Julie Margenthaler2, David J. Dooling4,5, David Ota23, Li-Wei Chang3,14, Ron Bose2,3, Timothy J. Ley1,2,4, David Piwnica-Worms8,9,10,12,24, Tammi L. Vickery4,5, Adnan Elhammali8,9,10, Helen Piwnica-Worms8,12,20,21, Sandra McDonald2,22, Mark Watson6,14,22, Michael C. Wendl4,5, Katherine DeSchryver1, D. Craig Allred3,14, Laura Esserman13,15, Gary Unzeitig16, Julie Margenthaler2, David J. Dooling4,5, David Ota23, Li-Wei Chang3,14, Ron Bose2,3, Timothy J. Ley1,2,4, David Piwnica-Worms8,9,10,12,24, Joshua M. Stuart11, Richard K. Wilson2,4,5 & Elaine R. Mardis2,4,5

To correlate the variable clinical features of oestrogen–receptor–positive breast cancer with somatic alterations, we studied pretreatment tumour biopsies accrued from patients in two studies of neoadjuvant aromatase inhibitor therapy by massively parallel sequencing and analysis. Eighteen significantly mutated genes were identified, including five genes (RUNXI, CBFβ, MYH9, ML3L and SF3B1) previously linked to haematopoietic disorders. Mutant MAP3K1 was associated with luminal A status, low-grade histology and low proliferation rates, whereas mutant TP53 was associated with the opposite pattern. Moreover, mutant GATA3 correlated with suppression of proliferation upon aromatase inhibitor treatment. Pathway analysis demonstrated that mutations in MAP2K4, a MAP3K1 substrate, produced similar perturbations as MAP3K1 loss. Distinct phenotypes in oestrogen–receptor–positive breast cancer are associated with specific patterns of somatic mutations that map into cellular pathways linked to tumour biology, but most recurrent mutations are relatively infrequent. Prospective clinical trials based on these findings will require comprehensive genome sequencing.

Oestrogen-receptor-positive breast cancer exhibits highly variable prognosis, histological growth patterns and treatment outcomes. Neoadjuvant aromatase inhibitor treatment trials provide an opportunity to document oestrogen-receptor-positive breast cancer phenotypes in a setting where sample acquisition is easy, prospective consent for genomic analysis can be obtained and responsiveness to oestrogen deprivation therapy is documented1. We therefore conducted massively parallel sequencing (MPS) on 77 samples accrued from two neoadjuvant aromatase inhibitor clinical trials2–5. Forty-six cases underwent whole-genome sequencing (WGS) and 31 cases underwent exome sequencing, followed by extensive analysis for somatic alterations and their association with aromatase inhibitor response. Case selection for discovery was based on the levels of the tumour proliferation marker Ki67 in the surgical specimen, because high cellular proliferation despite aromatase inhibitor treatment identifies poor prognosis tumours exhibiting oestrogen-independent growth1 (Supplementary Fig. 1). Twenty-nine samples had Ki67 levels above 10% (‘aromatase-inhibitor-resistant tumours’, median Ki67 21%, range 10.3–80%) and 48 were at or below 10% (‘aromatase-inhibitor-sensitive tumours’, median Ki67 1.2%, range 0–8%). Cases were also classified as luminal A or B by gene expression profiling1. We subsequently examined interactions between Ki67 biomarker change, histological categories, intrinsic subtype and mutation status in selected recurrently mutated genes in 310 cases overall. Pathway analysis was applied to contrast the signalling perturbations in aromatase-inhibitor-sensitive versus aromatase-inhibitor-resistant tumours.

Results
The mutation landscape of luminal–type breast cancer
Using paired-end MPS, 46 tumour and normal genomes were sequenced to at least 30-fold and 25-fold haploid coverage, respectively, with diploid coverage of at least 95% based on concordance with SNP array data (Supplementary Table 1). Candidate somatic events were identified using multiple algorithms4,6, and were then verified by hybridization capture-based validation that targeted all putative somatic single-nucleotide variants (SNVs) and small insertions/deletions (indels) that

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whole-genome sequencing of two pre-treatment biopsies was conducted. Digital sequencing data from captured target DNAs from the 46 tumour and normal pairs (Supplementary Table 2 and Supplementary Information) confirmed 81,858 mutations (point mutations and indels) and 773 somatic structural variants. The average numbers of somatic mutations and structural variants were 1,780 (range 44–11,619) and 16.8 (range 0–178) per case, respectively (Supplementary Table 3). Tier 1 point mutations and small indels predicted for all 46 cases were also validated using both 454 and Illumina sequencing (Supplementary Information). BRC25 was a clear outlier with only 44 validated tiers 1–3 mutations, all at low allele frequencies (ranging from 5% to 26.8%). This sample probably had low tumour content despite histopathology assessment, but the data are included to avoid bias.

The overall mutation rate was 1.18 validated mutations per megabase (Mb) (tier 1: 1.05; tier 2: 1.14; tier 3: 1.20). The mutation rate for tier 1 was higher than that observed for acute myeloid leukaemia (0.18–0.23)\(^6\), but lower than that reported for hepatocellular carcinoma (1.85)\(^9\), malignant melanoma (6.65)\(^9\) and lung cancers (3.05–8.93)\(^9\) (Supplementary Table 4). The background mutation rate (BMR) across the 21 aromatase-inhibitor-resistant tumours was 1.62 per Mb, nearly twice that of the 25 aromatase-inhibitor-sensitive tumours at 0.824 per Mb (\(P = 0.02\), one-sided \(t\)-test). A trend for more somatic structural variations in the aromatase-inhibitor-resistant group was also observed, as the validated somatic structural variation frequency in the 21 aromatase-inhibitor-resistant tumour genomes was 21.69 versus an average of 12.76 in 25 aromatase-inhibitor-sensitive tumours (\(P = 0.16\), one-sided \(t\)-test) (Fig. 1). If ten TP53 mutated cases were excluded, the background mutation rate still tended to be higher in the aromatase-inhibitor-resistant group (\(P = 0.08\)). To demonstrate that a single-tumour core biopsy produced representative genomic data, whole-genome sequencing of two pre-treatment biopsies was conducted for 5 of the 46 cases. The frequency of mutations in the paired specimens showed high concordance in all cases (correlation coefficient ranged from 0.74 to 0.95) (Supplementary Fig. 2) and a somatic mutation was infrequently detected in only one of the two samples (4.65% overall).

**Significantly mutated genes in luminal breast cancer**

The discovery effort was extended by studying 31 additional cases by exome sequencing, producing an additional 1,371 tier 1 mutations. In total the 77 cases yielded 3,355 tier 1 somatic mutations, including 3,208 point mutations, 1 dinucleotide mutation and 146 indels, ranging from 1 to 28 nucleotides. The point mutations included 733 silent, 2,145 missense, 178 nonsense, 6 read-through, 69 splice-site mutations and 77 in RNA genes (Supplementary Table 5). Of 2,145 missense mutations, 1,551 were predicted to be deleterious by SIFT\(^1\) and/or PolyPhen\(^2\). The MuSiC package\(^3\) was applied to determine the significance of the difference between observed versus expected mutation events in each gene, on the basis of the background mutation rate. This identified 18 significantly mutated genes with a convolution false discovery rate (FDR) <0.26 (Table 1 and Supplementary Table 6). The list contains genes previously identified as mutated in breast cancer (PIK3CA\(^4\), TP53\(^5\), GATA3\(^6\), CDH1\(^7\), RB1\(^8\), MLL3\(^9\), MAP3K1\(^10\) and CDKN1B\(^11\)) as well as genes not previously observed in clinical breast cancer samples, including TBX3, RUNX1, LDLRAPI, STNM2, MYH9, AGTR2, STMN2, SF3B1 and CBFB.

Thirteen mutations (3 nonsense, 6 frame-shift indels, 2 in-frame deletions and 2 missense) were identified in MAP3K1 (Table 1 and Fig. 2), a serine/threonine kinase that activates the ERK and JNK kinase pathways through phosphorylation of MAP2K1 and MAP2K4 (ref. 20). Of interest, a missense (S184L) and a splice-region mutation (e2+3 probably affecting splicing) in MAP3K1 were observed in two tumours with no MAP3K1 mutation (Fig. 2). Single nonsynonymous mutations in MAP3K12, MAP3K4, MAP4K3, MAP4K4, MAPK15 and MAPK3 were also detected (Supplementary Table 5). TBX3 harboured three small indels (one insertion and two deletions). TBX3 affects expansion of breast cancer stem-like cells through regulation of FGFR\(^21\). Two truncating mutations in the tumour suppressor CDKN1B were shared by the luminal A and B subsets. These were found in a total of 10 patients (3.9% of the series), a frequency close to that reported in a recent large-scale analysis of this gene in breast cancer (5.4%\(^22\)). The analysis supports a model in which CDKN1B acts as a tumour suppressor, with one copy of the gene sufficient to provide normal function. We found that CDKN1B is not mutated in normal breast tissue or matched normal breast tissue.

**Figure 1 | Genome-wide somatic mutations.** Circos plots\(^4\) indicate validated somatic mutations comprising tier 1 point mutations and indels, genome-wide copy number alterations, and structural rearrangements in six representative genomes. Three on-treatment Ki67 less than or at 10% (top panel: BRC15, BRC17 and BRC22) and three on-treatment Ki67 greater than 10% (bottom panel: BRC44, BRC47 and BRC50) cases are shown. Significantly mutated genes are highlighted in red. No purity-based copy number corrections were used for plotting copy number.
identified\(^*\). Four missense RUNX1 mutations were observed, with three in the RUNT domain clustered within the 8 amino acid putative ATP-binding site (R166Q, G168E and R169K). RUNX1 is a transcription factor affected by mutation and translocation in the M2 subtype of acute myeloid leukaemia\(^{27}\) and is implicated in tethering the oestrogen receptor to promoters independently of oestrogen response elements\(^{23}\). Two mutations (N104S and N140\(^{2}K\)) were also identified in CBFB, the binding partner of RUNX1. Additional mutations included 3 missense (2 K700E and 1 K666Q), in SF3B1, a splicing factor implicated in myelodysplasia\(^{4}\) and chronic lymphocytic leukaemia\(^{5}\). One missense mutation, one nonsense mutation and two indels were found in the MYH9 gene, involved in hereditary macrothrombocytopenia\(^{6}\) as well as being observed in an ALK translocation in anaplastic large cell lymphoma\(^{7}\).

We also identified three significantly mutated genes (LDLRAP1, AGTR2 and STM2N) not previously implicated in cancer. A missense and a nonsense mutation were observed in LDLRAP1, a gene associated with familial hypercholesterolaemia\(^{8}\). AGTR2, angiotensin II receptor type 2, harboured two missense mutations (V184I and R251H). Angiotensin signalling and oestrogen receptor interact in models of tissue fibrosis\(^{9}\). STM2N, a gene activated by JNK family kinases\(^{10,11}\) and therefore regulated by MAP3K1 and MAP2K4, harboured one frameshift deletion and one nonsense mutation. Three deletions and one point mutation (Supplementary Fig. 3) were identified in a large, infrequently spliced non-coding (inc) RNA gene, MALAT1 (metastasis associated lung adenocarcinoma transcript 1), that regulates alternative splicing by modulating the phosphorylation of SR splicing factor\(^{12}\). Translocations and point mutations of MALAT1 have been reported in sarcoma\(^{33}\) and colorectal cancer cell lines\(^{34}\). Five additional MALAT1 mutations were found in the recurrent screening set (Supplementary Table 5d). The locations of these mutations clustered in a region of species homology (F1 and 2 domains) that could mediate interactions with SRSF1 (ref. 32, Supplementary Fig. 4).

Non-coding mutation clusters were found in ATR, GPR126 and NRG3 (Supplementary Information and Supplementary Table 7).

### Correlating mutations with clinical data

To study clinical correlations, mutation recurrence screening was conducted on an additional 240 cases (Supplementary Table 8 and Supplementary Fig. 1). By combining WGS, exome and recurrence screening data, we determined the mutation frequency in PIK3CA to be 41.3% (131 of 317 tumours) (Supplementary Table 5a–d and Supplementary Fig. 3). TP53 was mutated in 51 of 317 tumours (16.1%) (Supplementary Table 5a–d and Supplementary Fig. 3). Additionally, 52 nonsynonymous MAP3K1 mutations in 39 tumours and 10 mutations in its substrate MAP2K4 were observed, representing a combined case frequency of 15.5% (Supplementary Table 5a–d and Fig. 3). Of note, 52 of the 62 non-silent mutations in MAP3K1 and MAP2K4 were scattered indels or other protein-truncating events strongly suggesting functional inactivation. In addition, 13 tumours harboured two non-silent MAP3K1 mutations, indicative of bi-allelic loss and reinforcing the conclusion that this gene is a tumour suppressor. Twenty nine tumours harboured a total of 30 mutations in GATA3, consisting of 25 truncation events, one in-frame insertion, and 4 missense mutations including 3 recurrent mutations at M294K (Supplementary Table 5a–d and Supplementary Fig. 3). BRC8 harboured a chromosome 10 deletion that includes GATA3. CDH1 mutation data were available for 169 samples and, as expected, its mutation status was strongly associated with lobular breast cancer (Table 2a). We applied a permutation-based approach in MuSiC\(^{16}\) to ascertain relationships between mutated genes. Negative correlations were found between mutations in gene pairs such as GATA3 and PIK3CA (\(P = 0.0026\), CDH1 and GATA3 (\(P = 0.015\)), and CDH1 and TP53 (\(P = 0.022\)). MAP3K1 and MAP2K4 mutations were mutually exclusive, albeit without reaching statistical significance (\(P = 0.3\)). In contrast, a positive correlation between MAP3K1/MAP2K4 and PIK3CA mutations was highly significant (\(P = 0.0002\)) (Supplementary Table 9).

Two independent mutation data sets, designated ‘Set 1’ (discovery cohort) and ‘Set 2’ (validation cohort), from these clinical trial samples were analysed separately and then in combination, with a false discovery rate (FDR)-corrected \(P\) value to gauge the overall strength and substitution, splice site mutation or indel is designated with a circle at the representative protein position with colour to indicate translation effects of the mutation. Asterisk, nonsense mutations that cause truncation of the open reading frame.

### Table 1 | Significantly mutated genes identified in 46 whole genomes and 31 exomes sequenced in luminal breast cancer patients

| Gene       | Total | MS | NS | Indel | SS | \(P\) value | FDR |
|------------|-------|----|----|-------|----|-------------|-----|
| MAP3K1     | 13    | 2  | 3  | 8     | 0  | 0           | 0   |
| PIK3CA     | 45    | 4  | 4  | 0     | 1  | 0           | 0.022|
| TP53       | 18    | 3  | 1  | 1     | 1  | 0.015       | 0.0022|
| GATA3      | 8     | 0  | 1  | 4     | 3  | 1.15 \(\times\) 10\(^{-19}\) | 7.41 \(\times\) 10\(^{-16}\) |
| CDH1       | 8     | 1  | 1  | 5     | 1  | 3.07 \(\times\) 10\(^{-15}\) | 1.59 \(\times\) 10\(^{-13}\) |
| TBX3       | 3     | 0  | 0  | 3     | 0  | 2.58 \(\times\) 10\(^{-6}\)  | 0.011|
| ATR        | 6     | 6  | 0  | 0     | 0  | 3.73 \(\times\) 10\(^{-6}\)  | 0.014|
| RUNX1      | 4     | 4  | 0  | 0     | 0  | 6.59 \(\times\) 10\(^{-6}\)  | 0.021|
| ENSG00000212670 | 2  | 2  | 0  | 0     | 0  | 2.31 \(\times\) 10\(^{-9}\)  | 0.066|
| RB1        | 4     | 2  | 1  | 0     | 0  | 2.76 \(\times\) 10\(^{-5}\)  | 0.071|
| LDLRAP1    | 2     | 1  | 1  | 0     | 0  | 4.27 \(\times\) 10\(^{-7}\)  | 0.092|
| STM2N      | 2     | 1  | 0  | 1     | 0  | 4.15 \(\times\) 10\(^{-6}\)  | 0.092|
| MYH9       | 4     | 1  | 1  | 2     | 0  | 8.96 \(\times\) 10\(^{-6}\)  | 0.178|
| MLL3       | 5     | 1  | 1  | 3     | 0  | 1.04 \(\times\) 10\(^{-6}\)  | 0.191|
| CBFB       | 2     | 0  | 1  | 1     | 0  | 1.39 \(\times\) 10\(^{-4}\)  | 0.240|
| AGTR2      | 2     | 2  | 0  | 0     | 0  | 1.71 \(\times\) 10\(^{-4}\)  | 0.256|
| SF3B1      | 3     | 3  | 0  | 0     | 0  | 1.79 \(\times\) 10\(^{-4}\)  | 0.256|
| STM2N      | 2     | 1  | 1  | 0     | 0  | 1.70 \(\times\) 10\(^{-4}\)  | 0.256|

*ENSG00000212670 is not in RefSeq release 50. MS, Missense; NS, nonsense; SS, splice site.

**Figure 2 |** MAP3K1 and MAP2K4 mutations observed in 317 samples. Somatic status of all mutations was obtained by Sanger sequencing of PCR products or Illumina sequencing of targeted capture products. The locations of conserved protein domains are highlighted. Each nonsynonymous mutation, splice site mutation or indel is designated with a circle at the representative protein position with colour to indicate translation effects of the mutation. Asterisk, nonsense mutations that cause truncation of the open reading frame.

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predictive marker for aromatase inhibitor response. This finding requires further verification because it was significant in sets of samples exhibiting greater percentage Ki67 decline. The mutation did not influence baseline Ki67 levels but was enriched in luminal B tumours when compared to the other three types. There were significantly more somatic NHEJ events in luminal B tumours than the other three types. Structural variant and DNA repair mechanisms

Analysis of copy number alterations (CNAs) revealed arm-level gains for 1q, 5q, 8q, 16q, 17q, 20p and 20q and arm-level losses for 1p, 8p, 16q, and 17p in the 46 WGS tumour genomes. A total of 773 structural variants (579 deletions, 189 translocations, and 65 inversions) identified using Illumina paired-end reads from whole-genome sequence data. One MAP3K1 deletion in BRC49 and one MAP2K4 deletion in BRC47, and one ELP3-NRG1 fusion in BRC49 identified using Illumina paired-end reads from whole-genome sequence data. Arcs represent multiple breakpoint-spanning read pairs with sequence coverage depth plotted in black across the region. Chr, chromosome.

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Pathways relevant to aromatase inhibitor response

Pathscan analysis (Supplementary Table 15 and Supplementary Information) indicated that somatic mutations detected in the 77 discovery cases affect a number of pathways, including caspase cascade/apoptosis, ErbB signalling, Akt/Pi3K/mTOR signalling, TP53/RB signalling and MAPK/JNK pathways. To discern the pathways relevant to aromatase inhibitor sensitivity, we conducted separate pathway analyses for aromatase-inhibitor-sensitive versus aromatase-inhibitor-resistant tumours. Whereas the majority of top altered pathways (FDR ≤ 0.15) in each group are shared, several pathways were enriched in the aromatase-inhibitor-resistant group, including the TP53 signalling pathway, DNA replication, and mismatch repair. Specifically, 38% of the aromatase-inhibitor-resistant group (11 of 29 tumours) have mutations in the TP53 pathway with three having double or triple hits involving TP53, ATR, APF1 or THBS1. In contrast, only 16.6% (8 of 48 tumours) of the Ki67 low group had mutations in the TP53 signalling pathway, each with only a single hit in genes TP53, ATR, CCNE2 or IGF1 (Supplementary Table 16).

GeneGo pathway analysis of MetaCore interacting network objects was used to identify genes in the 77 luminal breast cancers with low-frequency mutations that cluster into pathway maps. Eight networks assembled from significant maps encompassed mutations from 71 (92%) of the tumours (Fig. 4b). Many of the network objects shared pathways with significantly mutated genes such as TP53, MAP3K1, PIK3CA and CDH1. GeneGo analysis also revealed that several genes with low-frequency mutations were actually subunits of complexes, resulting in higher mutation rates for that object, for example, the condensin complex (4 mutations in 4 genes) and the MRN complex (4 mutations in 3 genes). Several pathways without multiple significantly mutated genes, such as the apoptotic cascade, calcium/phospholipase signalling and G-protein-coupled receptors, were significantly affected by low-frequency mutations. Grouping tumours by significantly mutated genes and pathway mutation status showed that whereas 55% (71 of 134) of the tumours contained significantly mutated genes in significant pathways, an additional 16% (21%) contained only non-significantly mutated genes in these pathways. Thus, tumours without a given significantly mutated gene often had other mutations in the same relevant pathway (Fig. 4b, Supplementary Fig. 6, Supplementary Table 17 and Supplementary Information).

We also applied PARADIGM to infer pathway-informed gene activities using gene expression and copy-number data to identify several ‘hubs’ of activity (Supplementary Fig. 7, Supplementary Fig. 8 and Supplementary Information). As expected, ESR1 and FOXA1 were among the hubs activated cohort-wide while other hubs exhibited high but differential changes in aromatase-inhibitor-resistant tumours including MYC, FOXM1 and MYB (Supplementary Fig. 8). The concordance among the 104 MetaCore maps from GeneGo analysis described above is significant, with 75% (72%) matching one of the
PARADIGM subnetworks at the 0.05 significance level after multiple test correction ($P < 4.4 \times 10^{-6}$, Bonferroni-adjusted hypergeometric test) (Supplementary Fig. 9). We identified significant subnetworks associated with Ki67 biomarker status (Supplementary Fig. 10 and Supplementary Information) involving transcription factors controlling large regulons.

The PARADIGM-inferred pathway signatures were further used to derive a map of the genetic mechanisms that may underlie treatment response. A subnetwork was constructed in which interactions were retained only if they connected two features with higher than average absolute association with Ki67 biomarker status (Supplementary Fig. 10 and Supplementary Information). Consistent with the PathScan results, among the largest of the hubs in the identified network were a central DNA damage hub with the second highest connectivity (55 regulatory interactions; 1% of the network) and TP53 with the 14th highest connectivity (26 connections; 0.5% of the network). Additional highly connected hubs identified in order of connectivity were MYC with 79 connections (1.4%), FYN with 45 (0.8%), MAPK3 with 43, JUN with 40, HDAC1 with 40, SHC1 with 39, and HIF1A/ARNT complex with 39 (Supplementary Fig. 11).

To identify higher-level connections between mutations and clinical features, we compared the samples on the basis of pathway-derived signatures. For each clinical attribute and each significantly mutated gene, we dichotomized the discovery samples into a positive and a negative group to derive pathway signatures that discriminated between the groups (see details in Supplementary Information). We then computed all pair-wise Pearson correlations between pathway signatures and clustered the resulting correlations (Fig. 5). The entire process was repeated using validated mutations and signatures derived from the validation set (Supplementary Fig. 12). In line with expectation, PIK3CA, MAP3K1, MAP2K4, and low risk preoperative endocrine prognostic index (PEPI) scores (PEPI is an index of recurrence risk post neoadjuvant aromatase inhibitor therapy) cluster with the luminal A subtypes and with each other, and are supported by the validation set analysis. The luminal B-like signatures included TP53, RB1, RUNX1 and MALAT1, which also associated over-represented by mutations in 77 luminal breast tumours (46 WGS and 31 exome cases). In the concentric circle diagram, tumours are arranged as radial spokes and categorized by their mutation status in each network (concentric ring colour) and significantly mutated gene mutation status (black dots). Tumour classification by pathway analysis shows many tumours unaffected by a given significantly mutated gene often harbour other mutations in the same network. For full annotation, see Supplementary Information and Supplementary Fig. 6. PLC, phospholipase C; SMG, significantly mutated gene.
with other poor outcome features such as high baseline and surgical Ki67 levels, high grade histology and high PEPI scores. The TP53 and MALAT1 associations in the discovery set also were supported by the validation set analysis.

**Druggable gene analysis**

We defined mutations in druggable tyrosine kinase domains including in ERBB2 (a V777L and a 755–759LRENT in-frame deletion homologous to gefitinib-sensitizing EGFR mutations in lung cancer37), as well as in DDR1 (A829V, R611C), DDR2 (E583D), CSF1R (D735H, M875L), and PDGFRA (E924K). In addition, plekstrin homology domain mutations were observed in AKT1 (C77F) and AKT2 (S11F) and a kinase domain mutation was identified in RPS6KB1 (S375F) (Supplementary Table 1).

**Discussion**

The low frequency of many significantly mutated genes presents an enormous challenge for correlative analysis, but several statistically significant patterns were identified, including the relationship between MAP3K1 mutation, luminal A subtype, low tumour grade and low Ki67 proliferation index. On this basis, for patients with MAP3K1 mutant luminal tumours, neoadjuvant aromatase inhibitor could provide a favourable option. In contrast, tumours with TP53 mutations, which are mostly aromatase inhibitor resistant, would be more appropriately treated with other modalities. MAP3K1 activates the ERK family, thus, loss of ERK signalling could explain the indolent nature of MAP3K1-deficient tumours20. However, MAP3K1 also activates JNK through MAP2K4, which also can be mutated38. Loss of JNK signalling produces a defect in apoptosis in response to stress, which would hypothetically explain why these mutations accumulate39,40. PIK3CA harboured the most mutations (41.3%) but was neither associated with clinical nor Ki67 response, confirming our earlier report41.

The positive association between MAP3K1/MAP2K4 mutations and PIK3CA mutation at both the mutation and pathway levels suggests cooperativity (Fig. 4a).

The finding of multiple significantly mutated genes linked previously to benign and malignant haematopoietic disorders suggests that breast cancer, like leukaemia, can be viewed as a stem-cell disorder.

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### Table 2 | Correlations between mutations and clinical features

**a** Luminal subtype and histology grade

| Gene   | Expression/histo-pathology variable | Mutation frequency* | Set1 *P† | Set2 *P† | Whole set FDR P‡ |
|--------|-------------------------------------|---------------------|----------|----------|------------------|
| TP53   | Luminal subtype A                   | 9.3% (13/140)       | 0.001    | 0.46     | 0.041            |
|        | Luminal subtype B                   | 21.5% (38/177)      |          |          |                  |
| TP53   | Histological grade I                | 4.5% (3/66)         | 0.05     | 0.067    | 0.02             |
| MAP3K1 | Luminal subtype A                   | 20.0% (28/140)      | 0.018    | 0.028    | 0.005            |
| MAP3K1 | Luminal subtype B                   | 6.2% (11/177)       |          |          |                  |
| CDH1   | Histological grade II/III           | 8.8% (22/250)       | 0.41‡    | 2.8 × 10⁻¹¹ | 3.9 × 10⁻¹⁰ |
|        | Histological type ductal            | 5.9% (10/169)       |          |          |                  |
|        | Histological type lobular           | 50.0% (20/40)       |          |          |                  |

**b** Mutation and Ki67 index

| Gene   | Ki67 variable          | Wild type mean | Mutant mean | Set1 *P* | Set2 *P* | Whole set FDR P* |
|--------|------------------------|----------------|-------------|----------|----------|------------------|
| TP53   | Baseline               | 13.1           | 25.1        | 3.7 × 10⁻⁵ | 0.012    | 0.0003           |
|        | Surgery                | 1.4            | 4           | 0.0002   | 0.014    | 0.001            |
|        | % change               | −92.4          | −84.3       | 0.09     | 0.28     | 0.24             |
| MAP3K1 | Baseline               | 15.8           | 8.1         | 0.049    | 0.001    | 0.002            |
|        | Surgery                | 1.86           | 0.75        | 0.11     | 0.1      | 0.05             |
|        | % change               | −88.3          | −90.5       | 0.49     | 0.65     | 0.55             |
| GATA3  | Baseline               | 14.8           | 11.5        | 0.13     | 0.95     | 0.56             |
|        | Surgery                | 1.95           | 0.38        | 0.001    | 0.23     | 0.012            |

* Unadjusted P value from Fisher’s exact test or Chi-square test as appropriate.
† Unadjusted P value from Fisher’s exact test or Chi-square test as appropriate.
‡ Only 77 cases in Set1 had CDH1 sequencing results.
§ Geometric means based on all cases (Set1 and Set2 combined).
* Unadjusted P value from Wilcoxon rank sum test.
that produces indolent or aggressive tumours that display varying phenotypes depending on differentiation blocks generated by different mutation repertoires.\textsuperscript{2} Whereas only MLL3 showed statistical significance in the analysis of 46 WGS cases, multiple mutations in genes related to histone modification and chromatin remodelling are worth noting (Supplementary Table 19). An array of coding mutations and structural variations was discovered in methyltransferases (MLL2, MLL3, MLL4 and MLL5), demethyltransferases (KDM6A, KDM4A, KDM5B and KDM5C), and acetyltransferases (MYST1, MYST3 and MYST4). Furthermore, we identified several adenine-thymine (AT)-rich interactive domain-containing protein genes (ARID1A, ARID2, ARID3B and ARID4B) that harboured mutations and large deletions, reinforcing the role of members from the SNF/SWI family in breast cancer.

Pathway analysis enables the evaluation of mutations with low recurrence frequency where statistical comparisons are conventionally underpowered. For example, the eight samples with MAP2K4 mutations were sufficient to derive a reliable pathway-based gene signature in PARADIGM that aligns with MAP3K1. This approach also pointed to a putative connection between MALAT1 and the TP53 pathway. Finally, we provide evidence that transcriptional associations to Ki67 response reside in a connected network under the control of several key ‘hub’ genes including MYC, PYN and MAP kinases, among others. Targeting these hubs in resistant tumours could produce therapeutic advances. In conclusion, the genomic information derived from unbiased sequencing is a logical new starting point for clinical investigation, where the mutation status of an individual patient is determined in advance and treatment decisions are driven by therapeutic hypotheses that stem from knowledge of the genomic sequence and its possible consequences. However, the accrual of large numbers of patients and the use of comprehensive sequencing and gene expression approaches will be required because of the extreme genetic heterogeneity documented by this investigation.

METHODS SUMMARY

Clinical trial samples were accessed from the preoperative letrozole phase 2 study (NCT00084396\textsuperscript{2}) that investigated the effect of letrozole for 16 to 24 weeks on surgical outcomes and from the American College of Surgeons Oncology Group (http://www.genego.com/metacore.php) and PARADIGM. A complete description of the materials and methods used to generate this data set and results is provided in the Supplementary Methods section.

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Author Contributions M.J.E. led the clinical investigations, biomarker analysis and chip-based genomics. E.R.M., M.J.E., L.D., R.S.F., T.J.L. and R.K.W. designed the experiments. L.D. and M.J.E. led data analysis. D.S., J.W.W., D.C.K., C.C.H., M.D.M., K.C., C.A.M., F.D., W.S.S., M.C.W., R.C. and C.K. performed data analysis. D.S., C.A.M., J.W.W., J.F.M., C.L. and L.D. prepared figures and tables. R.S.F., L.L.F., R.D., M.H., T.L.V., J.H., L.L., R.C. and J.S. performed laboratory experiments. L.E., G.U., J.M., G.V.B., P.K.M., J.M.G., M.L., K.H. and J.O. provided samples and clinical data. V.J.S., K.B., J.L., Y.T. and C.K. provided statistical and clinical correlation analysis. D.O. oversees the ACOSOG Operations Center that provides oversight and tracking for ACOSOG clinical trials. K.D., S.Mc.D., D.C.A. and M.W. provided pathology analysis. B.A.V.T., J.W., R.J.G., A.E., D.P.-W., H.P.-W., J.M.S., T.C.G., S.N., C.K. and M.C.W. performed pathway analysis. L.-W.C. and R.B. analysed the druggable target mutation data. D.J.D. and B.O. provided informatics support. L.D., M.J.E. and E.R.M. wrote the manuscript. T.J.L., M.C.W. and R.K.W. critically read and commented on the manuscript.

Author Information DNA sequence data are deposited in the restricted access portal at dbGaP, accession number phs000472.v1.p1. Gene expression array data used in the Paradigm training set is deposited in GEO, accession number GSE29442, and a Superset that covers both the Agilent gene expression data and the Agilent array CGH data used for the Paradigm test set is deposited in GEO, accession number GSE35191. Reprints and permissions information is available at www.nature.com/reprints. This paper is distributed under the terms of the Creative Commons Attribution-Non-Commercial-Share Alike licence, and is freely available to all readers at www.nature.com/nature. The authors declare no competing financial interests. Readers are welcome to comment on the online version of this article at www.nature.com/nature. Correspondence and requests for materials should be addressed to M.J.E., L.D. and E.R.M.