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# Gene Expression Meta-Analysis identifies Cytokine Pathways and 5q Aberrations involved in Metastasis of ERBB2 Amplified and Basal Breast Cancer

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**Background:** Breast tumors have been described by molecular subtypes characterized by pervasively different gene expression profiles. The subtypes are associated with different clinical parameters and origin of precursor cells. However, the biological pathways and chromosomal aberrations that differ between the subgroups are less well characterized. The molecular subtypes are associated with different risk of metastatic recurrence of the disease. Nevertheless, the performance of these overall patterns to predict outcome is far from optimal, suggesting that biological mechanisms that extend beyond the subgroups impact metastasis.

**Results:** We have scrutinized publicly available gene expression datasets and identified molecular subtypes in 1,394 breast tumors with outcome data. By analysis of chromosomal regions and pathways using "Gene set enrichment analysis" followed by a meta-analysis, we identified comprehensive mechanistic differences between the subgroups. Furthermore, the same approach was used to investigate mechanisms related to metastasis within the subgroups. A striking finding is that the molecular subtypes account for the majority of biological mechanisms associated with metastasis. However, some mechanisms, aside from the subtypes, were identified in a training set of 1,239 tumors and confirmed by survival analysis in two independent validation datasets from the same type of platform and consisting of very comparable node-negative patients that did not receive adjuvant medical therapy. The results show that high expression of 5q14 genes and low levels of TNFR2 pathway genes were associated with poor survival in basal-like cancers. Furthermore, low expression of 5q33 genes and interleukin-12 pathway genes were associated with poor outcome exclusively in ERBB2-like tumors.

Conclusion: The identified regions, genes, and pathways may be potential drug targets in future individualized treatment strategies.

Keywords: breast cancer, metastasis, gene expression, microarray, pathway analysis, molecular subtypes

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## Background

Breast cancer is the most common cancer among women. The local disease is not lethal, but its spread to distant organs is fatal. The risk of metastasis is evaluated by clinical and pathological criteria, but the performance of these methods is far from optimal. Gene expression profiling of tumors has been used for the supervised classification of cancer outcomes in several studies with promising results for the improvement of risk prediction.<sup>1-4</sup> Despite these promising clinical results, limited insights into its biological mechanisms has been obtained from the large amount of gene expression data available. A very different approach was used in early studies of global gene expression of breast cancer, where unsupervised hierarchical clustering was used to identify 4–5 molecular subtypes.<sup>5–8</sup> These subtypes arise from at least two cell types: basal cells and luminal epithelial cells. Luminal tumors are mainly estrogen receptor (ER)-positive and cluster in two distinct groups (termed luminal A and luminal B) that appear to result in good and poor prognosis, respectively. Basal-like tumors are ER-negative and are characterized by poor prognosis. A distinct group that mainly consists of ER-negative tumors is characterized by the amplification of ERBB2. Finally, a normal-like profile constitutes good prognosis. These subtypes are biologically very meaningful, but their prediction of metastasis is not optimal.

In recent studies, unsupervised and supervised methods have been used to differentiate the molecular subtypes of breast cancer into further subgroups with different outcomes. The discriminating genes have subsequently been related to biological function (for example, immune response among ERnegative tumors).9,10 These studies have identified very broad mechanisms, but they have not pointed at specific biochemical pathways. A very different strategy would be to analyze predefined gene sets representing biological mechanisms, such as the pathways for their association to metastasis. We have previously used this strategy for the analysis of the overall pathways involved in the metastasis of breast cancer.<sup>11</sup> Furthermore, we have applied this approach to gene sets representing genomic regions to identify somatic mutations at the deoxyribonucleic acid (DNA)-level that are involved in metastasis.<sup>12</sup> However, the analysis of pathways and genomic aberrations has not been applied to analyze



the various metastatic mechanisms found within the different subtypes of breast cancer.

We hypothesize that several somatic genomic aberrations and biological pathways characterize the molecular subtypes in breast cancer. Despite the association of the subtypes to metastasis, we expect that mechanisms within a given subtype may be involved in the higher risk of metastasis for a fraction of the tumors. Furthermore, we anticipate that deregulation of the pathways is reflected in gene expression patterns of primary tumor biopsies. We also assume that somatic copy number mutations involving chromosomal regions will be reflected in the overall level of gene expression in these regions. Our aim is to elucidate these metastatic mechanisms within the molecular subtypes and, secondly, to determine the mechanisms that differ between the subgroups.

We have used predefined gene sets representing canonical pathways and chromosomal regions to identify metastatic mechanisms within molecular subtypes. This has been accomplished by a global approach; gene set enrichment analysis (GSEA) examining the entire list of genes ranked according to association to metastasis within each molecular subtype. In a single dataset, this analysis will generally not result in significant findings. However, by performing a meta-analysis of several datasets, we have increased the power to identify somatic mutations among the chromosomal regions and pathways involved in the metastasis of fractions of breast tumors within a specific subtype.

#### Results

#### Molecular subtypes of 1,439 tumors

Global gene expression data were collected from 1,586 breast tumors with clinical follow up (Table 1). Molecular subtypes were identified by a single sample prediction approach using nearest centroid classification.<sup>8</sup> Single-sample predictions resulted in 1,394 samples with assigned subtypes. Survival analysis confirmed the prognostic impact of the subtypes, with luminal A and normal-like tumors having significantly better prognosis than luminal B, ERBB2, and basal-like tumors (Fig. 1).

# Genomic aberrations and pathways associated to molecular subtypes

To identify the biological characteristics of the molecular subtypes, we divided the 1,394 samples



Data set	Country	Platform	#genes	Outcome	#pt with follow up	#sub typed samples
Amsterdam <sup>1</sup>	The Nederlands	Agilent	24451	Distant metastasis	295	287
Rotterdam <sup>2</sup>	The Nederlands	133A	22283	Distant metastasis	286	277
Stockholm <sup>32</sup>	Sweden	133A+B	44982	Death from BC	159	152
Uppsala <sup>33</sup>	Sweden	133A+B	44982	Death from BC	236	229
Bild <sup>34</sup>	US	95av2	12625	Metastasis	158	153
Hu <sup>8</sup>		3 Agilent	22K	Relapse	105	105
Training total		Ũ		·	1239	1203
Mainz <sup>9</sup>	Germany	133A	22283	Distant metastasis	200	191
TRANSBIG-S <sup>3</sup>	UK	133A	22283	Distant metastasis	147	0
Test total					347	191
Total					1586	1394

Table 1. Datasets and	patients	included	in the	e study.
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subtyped with a single sample prediction, to a training set of 1,203 tumors corresponding to six entire datasets and a test set of 191 tumors contained in two datasets (Table 1). We applied GSEA analysis to one subtype at a time compared to all other tumors. This analysis was performed with gene sets representing chromosomal region and with pathway gene sets. A meta-analysis was performed to identify gene sets that were concordantly differentially regulated in the training datasets. For chromosomal regions, a large number of somatic mutations were indicated by the analysis (Table 2). The differential expression of several consecutive regions indicated that larger regions are lost or gained. Validation in the test set resulted in



Figure 1. Kaplan–Meyer plot of the five molecular subtypes in seven datasets, including 1,394 tumor samples. Molecular subtypes were determined by the single-sample prediction method reported by Hu et al.<sup>8</sup> Hazard ratios were calculated by fitting a Cox model to the data for comparing each group to basal-like tumors. The *P*-value was calculated using the log-rank test.



Basal	ERBB2	Lum A	Lum B	Normal
Up-regulated				
1P34, 1P32, 1P22	2P11, 2Q37	1P33, 1P31, 1Q41	4P16	2Q32
2P24, 2P21, <b>2P16</b>	6Q21	3P22, 3P21, 3P14	6P22	3P22
3Q21, <b>3Q25</b> , <b>3Q27</b> , 3Q29	9Q34	5Q11, 5Q13	8P11, <b>8Q22</b> , <b>8Q24</b>	<b>5Q13</b> , 5Q32
6P21, 6Q23	12Q12	8P22, 8P21	<b>12Q13</b> , 12Q24	7P15
<b>7P15</b> , 7Q32	16Q13	14Q24	16P13, 16P12, 16P11	8P23, 8P21
10P15	<b>17Q11</b> , 17Q12,	10Q23, 10Q25	<b>17Q21</b> , 17Q22, 17Q23,	9Q21
11Q22	17Q21, <b>17Q23</b>	17P11	17Q24, 17Q25	11Q23
12P13, 12P12	22Q11, 22Q13		20P11, <b>20Q11</b> , <b>20Q12</b> ,	15Q11
16Q13, <b>16Q24</b>			20Q13	XQ13
21Q22				XQ22
XQ26, XQ28				
Down-regulated				
3P22, 3P21, 3P14	1P21	2P25, 2P12	<b>1P31</b> , 1P22	1Q23, 1Q41, <b>1Q42</b>
4P16	3P24	3Q27	2Q32	2P12
<b>5Q11</b> , <b>5Q13</b> , 5Q14, 5Q31	8P22, 8P21	6P21	4P15, 4Q21	6P22
9Q32	9P24	8Q22, 8Q24	5P13	8P12, 8Q22, 8Q24
10Q23, <b>10Q25</b>	17P11	10P15	6P25, 6Q21, <b>6Q22</b> , <b>6Q23</b>	10Q21
12Q12, <b>12Q13</b> , 12Q22	18Q12, 18Q21	16Q13, <b>16Q22</b> , <b>16Q24</b>	<b>7P15</b> , 7Q31	17Q25
14Q21, <b>14Q23</b> , 14Q24,		17Q11, <b>17Q23</b> , 17Q25	9P24, 9Q21	20Q13
14Q32		20P13, 20Q11, <b>20Q13</b>	<b>11Q22</b> , 11Q23, <b>11Q24</b>	
15Q22		22Q11, 22Q13		
16P13, 16P12		XQ28		
17P11, 17Q21, 17Q24				

Table 2. Differentially expressed chromosomal regions.

Note: Gene sets in bold were also significant in test data set (Mainz).

34% of regions being significant (highlighted in bold in Table 2).

Pathway analysis using predefined canonical pathways also indicated large differences between the subgroups (Table 3). Of these pathways, 43% were also identified by GSEA analysis in the independent Mainz dataset (highlighted in bold in Table 3).

# Chromosomal regions and pathways involved in metastasis within subgroups

The above analyses indicate that molecular subtypes significantly account for the metastasis of breast cancer, and that the subtypes are associated with very different biological pathways and somatic mutations. To identify the metastatic mechanisms that extend beyond the molecular subtypes, we performed pathway and chromosomal aberration analyses within each subtype, comparing tumors from patients that experienced metastasis to tumors from patients that did not experience metastasis within the follow-up period. The data were again split into a training set including six datasets, and the two most recently published datasets (Mainz and TRANSBIG-S) were kept for validation purposes. The training set was composed of datasets with different clinical characteristics and treatment statuses of the patients and performed with different microarray platforms. In contrast, the test sets (Mainz and TRANSBIG-S) contained data from node-negative patients not receiving adjuvant medical treatment; all analyses were performed with Affymetrix HG133A chips. The GSEA meta-analysis in the training set resulted in the identification of nine significant regions and pathways (Table S1). Using GSEA for the validation of these regions and pathways in the Mainz dataset did not result in any significant gene sets, probably due to the limited sample size when comparing tumors with poor and good outcomes within subtypes (data not shown).

In order to validate the identified regions and pathways with a different method, we investigated whether the overall expression level in each gene set could be used as a prognostic marker in survival analysis. This would also allow us to compare the expression of regions or pathways between single tumors from all datasets. Furthermore, utilizing survival information might help to improve the power of the validation, and combining datasets would also increase the power. Datasets were preprocessed and combined

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Table 3 Differentially expre-	ssed pathway gene sets.			
Basal	ERBB2	Lum A	Lum B	Normal
Up-regulated CELL CYCLE(6) DNA_REPLICATION(3) tRNA SYNTHESIS(2) ACTINY TNF APOPTOSIS(2) ACTINY TNF APOPTOSIS(2) CELL_ADHESION PROTEASOME ATRBRCA PS3 SIGNALING IL12(2) TH1TH2 S3 SIGNALING IL12(2) TH1TH2 ST_B_CELL_SIGNAL TRANSDUCTION ANTIGEN_RECEPTOR ST_B_CELL_CYTOTON ANTIGEN_RECEPTOR ST_FAS_SIGNALING CIPCOSPHINGOLIPID_ BIOSYNTH ST_FAS_SIGNALING ST_FAS_SIGNALING ST_FAS_SIGNALING Down-regulated	BIOSYNTHESIS_OF_ STEROIDS(2) PROTEASOME(2) ARGININE/PROLINE_ METABOLISM	AMINO ACID METABOLISM(4) ST_JNK_MAPK CIRCADIAN_EXERCISE NITROGEN_ METABOLISM(2) BUTANOATE METABOLISM(2) PROPANOATE METABOLISM ESTROGEN SIGNALING SPHINGOLIPID METABOLISM FATTY_ACID METABOLISM FATTY_ACID METABOLISM AKT IGF1 GLEVEC CARM_ER PDGF	OXIDATIVE PHOSPHORYLATION(3) MPR P53 P53 P53 P53 P53 P53 P50 PTOIN METABOLISM(2) PTDIN PTDIN DNA_REPLICATION(3) CELL_CYCLE TRANSCRIPTION CELL_CYCLE TRANSCRIPTION RAS CARM_ER CARM_CARD BIOSYNTHESIS_OF_ STEROIDS CCRM_CARD BIOSYNTHESIS_OF_ STEROIDS CCRM_CARD BIOSYNTHESIS_OF_ CCREB CCREB CCREB BIOSYNTHESIS_OF_ CCREB	FATTY_ACID_METABOLISM HISTIDINE_METABOLISM MAPK_SIGNALING WNT_SIGNALING WNT_SIGNALING(3) FOCAL_ADHESION ECM PROSTAGLANDIN CARDIACEG ALK PDGF ALK PDGF PDGF PDGF PDGF PDGF PDGF PDGF PDGF
TEL MTOR(2) IGF1(2) GPCR VIP AMINO ACID METABOLISM(2) ESTROGEN SIGNALING NUCLEAR SIGNALING NUCLEAR SIGNALING NUCLEAR SIGNALING CERAMIDE CERAMIDE CERAMIDE CERAMIDE CERAMIDE CERAMIDE CERAMIDE CERAMIDE CERAMIDE CERAMIDE CERAMIDE CERAMIDE CERAMIDE ST_JNK_MAPK BAD	<b>TEL</b> GLEEVEC IGF1 NITROGEN_ METABOLISM(2)	CELL_CYCLE(6) IL12(2) GLYCOLYSIS FENTOSE PENTOSE PHOSPHATE_(2) PROTEASOME ST_B_CELL_ANTIGEN_ RECEPTOR PTDINS DNA_REPLICATION(3) tRNA_SYNTHESIS(2) ST_T_CELL_SIGNAL TRANSDUCTION ACTINY CELL_ADHESION S3_SIGNALING NATURAL_KILLER_ CELL_CYTOTOX	PROSTAGLANDIN_ SYNTHESIS NTHI ACTIN_ CYTOSKELETON GLYCEROLIPID_ METABOLISM ILIR TOLL LYSINE_DEGRADATION MELANOGENESIS FOCAL_ADHESION WNT_SIGNALING ALK CELL_ADHESION(2) CYTOKINE UREA_CYCLE	CELL_CYCLE(5) DNA_REPLICATION(5) PENTOSE_PHOSPHATE(2) TRNA_BIOSYNTHESIS(2) OXIDATIVE_ PHOSPHORYLATION(3) CITRATE_CYCLE(3) PROTEASOME(3) GLYCOLYSIS PS3 BIOSYNTHESIS_OF_STEROIDS(2) ATRBRCA ST_FAS_SIGNALING GLUTAMATE_METABOLISM SELENOAMINO_ACID_ MPR RACCYCD MPR
Notes: Gene sets in bold were als	so significant in test data set (Mai	nz). The number of redundant pathwa	ay gene sets is given in brackets.	

using only genes measured on all chips. The expression level of all genes in each region or pathway identified previously was implemented in the region scores. This method is different from the GSEA metaanalysis described earlier because only genes represented on all platforms were included in the analysis. In 280 basal-like tumors from the training sets, this different approach supported a high level of expression of 5q14 (Fig. 2A) and a low level of expression of tumor necrosis factor receptor (TNFR)2 (Fig. 2E) as predictors of poor outcomes. For ERBB2 tumors, the low expression of 5q33 (borderline significant; Fig. 3A) and interleukin (IL)-12 (Fig. 3E) were associated with poor outcomes, which was in agreement with the GSEA meta-analysis. The remaining five pathways and regions identified by the GSEA metaanalysis were not significant in the survival analysis (Table S1).

For the validation of metastatic gene sets in independent datasets, two test sets were used. The Mainz data set, where 41 basal-like tumors were identified by single-sample prediction, was used for the validation of the metastatic mechanisms of basal-like tumors. However, only 20 ERBB2 tumors were identified by single-sample prediction in the Mainz dataset. To increase the sample size of ERBB2-positive tumors, the TRANSBIG-S dataset was included. The identification of molecular subtypes failed in this dataset, but ERBB2 status was determined from ERBB2 expression in both datasets. The very unbalanced distribution of ERBB2 and the coregulation of GRB7 located closely to ERBB2 supported this method for the classification of ERBB2 amplification (Fig. S1).

For basal-like tumors, the disadvantage of high 5q14 expression was validated by the survival analysis in the test set (Fig. 2C). To investigate whether this prognostic effect of 5q14 aberration was specific for basal-like tumors, the prognostic performance was also examined in the remaining tumors not belonging to this subgroup. This demonstrated no prognostic impact of 5q14 aberrations in the training set (Fig. 2B) but, surprisingly, a borderline significant opposite effect was found in the test set (Fig. 2D).

The prognostic impact of the TNFR2 pathway identified in the GSEA meta-analysis and in the survival analysis in the training set (Fig. 2E) was also supported by a survival analysis in the test set (Fig. 2G). However, this pathway was also significant in nonbasal-like tumors in the training set, but with smaller separation of the groups (ie, lower hazard ratios [Fig. 2F], and the pathway was only borderline significant among the nonbasal-like tumors in the test set [Fig. 2H]; in addition, a lower hazard ratio was observed among nonbasal-like tumors when compared to the basal-like tumors).

For ERBB2 tumors, the low expression of 5q33 (Fig. 3C) and IL-12 (Fig. 3G) was also associated with poor outcomes in the test set. Interestingly, these mechanisms were specific for tumors with high levels of expression of ERBB2 (Fig. 3B, D, F, and H).

To identify single metastasis candidate genes in the two 5q regions, we compared the expression of metastasizing and non-metastasizing tumors at single gene level. For 5q14, one gene, *HAPLN1*, was strongly and differentially regulated (Fig. S2), suggesting that this is a plausibly causal metastasis gene. No single gene was strongly and differentially expressed at 5q33 (data not shown).

#### Discussion

We have conducted a meta-analysis of the tumor gene expression data and identified chromosomal regions and pathways that were strongly associated with the molecular subtypes of breast cancer. Furthermore, some mechanisms were found to be associated with metastasis within the different subgroups. Our results demonstrate that these mechanisms are reflected in the overall gene expression of the chromosomal region and the pathway gene sets, which is in agreement with our hypotheses.

A major finding is the large number of chromosomal regions and pathways associated with the molecular subtypes that were found. This is in agreement with the supposedly different cellular origins of basallike and luminal-like tumors. The universal nature of the subtypes is supported by several studies9,10 that have used different unsupervised methods to identify the subtypes of breast tumors that resemble the subgroups reported by Sørlie et al.<sup>6</sup> ER status, ERBB2 status, and proliferation level are believed to be major characteristics of the molecular subtypes. Our results support the crucial impact of the following mechanisms: 17q amplification in ERBB2-like tumors (Table 2); estrogen signaling in luminal cancers (CARM ER and ESTROGEN SIGNALING; Table 3); and elevated cell cycle in basal-like and





Figure 2. Kaplan–Meyer plots of the 5q14 and TNFR2 groups in basal-like tumors. The training set consisted of 280 basal-like tumors (**A** and **E**) and 959 nonbasal-like tumors (**B** and **F**). The test set contained 41 basal-like tumors (**C** and **G**) and 159 nonbasal-like tumors (**D** and **H**). The score is calculated for each sample as the mean standardized expression values for all genes in the gene set The low score group was defined as the third of samples with the lowest score for the indicated gene set, and the high score group was defined as the third of the samples with the highest score. The middle third was excluded from the analysis. The *P*-values were calculated using the log-rank test. **Abbreviation:** TNFR, tumor necrosis factor receptor.





**Figure 3.** Kaplan–Meyer plots of 5q33 and IL-12 groups in ERBB2 tumors. The training set consisted of 196 tumors with high ERBB2 expression (**A** and **E**) and 1,043 tumors with low ERBB2 expression (**B** and **F**). The test set contained 70 tumors with the highest ERBB2 expression (**C** and **G**) and 277 tumors with lowest ERBB2 expression (**D** and **H**). The score is calculated for each sample as the mean standardized expression values for all genes in the gene set. The low score group was defined as the third of samples with lowest score for the indicated gene set, and the high score group was defined as the third of the samples with the highest score. The middle third was excluded from the analysis. The *P*-values were calculated using the log-rank test. **Abbreviation:** IL, interleukin.



luminal B tumors. By comparing metastasizing and nonmetastasizing tumors, without stratifying for subtypes, we have previously identified the chromosomal regions and pathways involved in the metastasis of breast cancer.<sup>11,12</sup> The identified gene sets strongly overlap with the gene sets that were differentially expressed between molecular subtypes, indicating that the molecular subtypes account for a majority of the metastasis-associated mechanisms. This is strongly supported by the very few metastasis-linked mechanisms we have identified within the molecular subgroups (two regions and two pathways), even though the cut-off for the false discovery rate (FDR) was 0.05 for intra-subgroup analysis and 0.01 for the between-subgroup analysis. However, the considerably smaller sample size in the within-group analysis may bias this conclusion.

Our within-subgroups analysis identified somatic mutations, which were located at different loci on 5q, and which had opposite prognostic effects: high levels of expression of 5q14 were associated with poor outcomes in basal-like cancer, whereas lower levels of expression of 5q33 were associated with poor outcomes among patients with ERBB2-amplified breast tumors. The dual role of 5q aberrations is supported by a few previous publications that used comparative genome hybridization; Friedrich et al<sup>13</sup> reported higher frequency of 5q23 losses among cases with distant metastasis and Karlsson et al<sup>14</sup> showed that poorer outcomes were associated with 5q31~qter gain for cases with ER-negative tumors. Callaghan et al<sup>15</sup> reported a higher frequency of allelic imbalance at 5q21 in patients with lymph node metastasis, indirectly supporting the prognostic significance of this region. However, these studies included few patients and did not subdivide tumors according to basal/ ERBB2 status, hampering the conclusion for these aberrations that, according to our results, are specific for basal-like and ERBB2 tumors, respectively.

Our method identifies regions by measuring the differential expression of genes, but it does not (in principle) determine whether the DNA copy number increases in one outcome group or decreases in the opposite group. However, low expression of 5q14 and its surrounding region characterize basal-like tumors according to our analysis, which compared molecular subtypes (Table 2). This is also supported by Adélaïde et al,<sup>16</sup> who used array comparative

genomic hybridization (aCGH) and reported 5q loss to be frequent in basal-like tumors. This indicates that the loss of 5q14 is common in basal-like tumors, but that patients with basal-like tumors retaining 5q14 have a worse prognosis. This might be explained by the presence of a tumor suppressor gene involved in basal-like cancer development, and another gene promoting metastasis-both of which were located in the region. A similar effect has previously been observed for 16q loss in breast cancer.<sup>12,17</sup> We pinpointed one gene, hyaluronan, and proteoglycan link protein 1 (HAPLN1) in the 5q14 region, which had a strong association with the metastasis of basal-like cancers, making this a relevant candidate gene that may be a causal factor in the metastasis of basal-like tumors. HAPLN1 is involved in the stabilization of hyaluronan and vesican. An interaction of hyaluronan and its receptor (CD44) induces signaling events that promote cell growth, migration, and metastasis,<sup>18</sup> making genes in this pathway relevant.

ERBB2-like tumors are not characterized by 5q aberrations, as compared to other subtypes (Table 2), but our results indicate that low expression of the 5q33 region is associated with poor outcomes. This may be the result of 5q33 loss in the poor prognosis group, or gain of the region in the good prognosis group. However, this cannot be determined from the present results.

Many pathways have previously been linked to metastasis; however, investigations of the metastatic pathways within different subtypes of breast cancer are sparse, as will be discussed in the following. We identified the high expression of the TNFR2 pathway as an indicator of good prognosis in basal-like cancers. TNFR2 is a receptor with equal affinity for tumor necrosis factor (TNF)a and TNFb. Both of these are produced by activated lymphocytes and can be toxic to many tumor types. Nevertheless, for micropapillary breast carcinomas, overexpression of TNFR2 has been linked with higher incidence of lymph node metastasis, which may conflict with our results.<sup>19</sup> However, our finding that the TNFR2 metagene is associated with survival in basal-like and nonbasal-like cancers in both the training set and test set (borderline significance for the nonbasal-like tumors in the test set) strongly indicate the prognostic disadvantage of low TNFR2 pathway expression. According to our results (Table 3), the TNFR2 pathway is not differentially

expressed between subgroups. However, the survival analysis within subtypes show smaller separations of high and low TNFR2 score curves (ie, lower hazard ratios for nonbasal-like tumors compared to basallike tumors; Fig. 2), indicating stronger metastatic prevention by the pathway in basal-like tumors. This is also supported by the GSEA meta-analysis that exclusively identified TNFR2 only among basal-like carcinomas. To our knowledge, no previous studies have described a prognostic influence of TNFR2/ TNF signaling among basal-like breast tumors.

The other pathway, IL-12, identified within the ERBB2 subgroup is also associated with good outcomes. This is supported by studies showing that IL-12 activates natural killer cells and enhances their ability to kill tumor cell lines that are treated with herceptin<sup>20</sup> However, no benefit was observed by adding IL-12 to the herceptin treatment of breast cancer patients in a small Phase 1 study.<sup>21</sup> None of the patients in our analysis received herceptin, indicating that IL-12 signaling has antimetastatic potential, independent of herceptin treatment.

Both IL-12 and TNFR2 signaling are involved in immune activation, suggesting that lymphocyte infiltration is beneficial for patients with ER-negative tumors. This is supported by pathological investigations conducted among subgroups of patients. For instance, Ménard et al<sup>22</sup> reported selectively higher survival rates in young breast cancer patients when they had lymphocyte infiltrated tumors, and Aaltomaa et al<sup>23</sup> described the benefit of lymphocyte infiltration among patients with rapid proliferating tumors. Several recent publications using gene expression datasets have also identified immune-related gene signatures that distinguish groups with different outcomes. Schmidt et al9 used hierarchical clustering to identify a B-cell signature with a prognostic impact among high-proliferating tumors. Teschendorff et al<sup>10</sup> focused their analysis on ER-negative tumors and used an unsupervised clustering technique to identify five subclusters. Among tumors with normal ERBB2 status, they found an immune response gene cluster that was associated with survival. Alexe et al<sup>24</sup> also used unsupervised clustering of ERBB2-positive tumors and reported a subtype with upregulated lymphocyte infiltration signal and improved survival. Rody et al<sup>25</sup> clustered immune-related genes and identified seven different gene clusters that were shown to represent

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Our comparison of molecular subtypes has identified comprehensive differences in pathways and genomic aberrations (Tables 2 and 3). Several of the somatic aberrations are supported by previous studies using aCGH to compare relevant groups. Bergamaschi et al<sup>27</sup> examined 89 tumors by gene expression profiling to identify four molecular subtypes; Adélaïde et al<sup>16</sup> compared 44 basal and 49 luminal tumors via aCGH, and a recent large study<sup>28</sup> performed a metaanalysis of 773 tumors. The results from these studies confirmed many of the regions identified in the pres-





ent investigation. For basal-like tumors, their results supported the extensive loss of 5q, as well as the loss of 10q23, 12q22, 14q21, 14q23, 15q22, and the gain of 3q25-27, 6p21, 7q32, 10p15, 12p13-12, and 21q22. For luminal A tumors, they also observed 1q41 and 8p22-21 gains and a 16q loss. For luminal B tumors, gains of 8q22-24, 20p11, and 20q13 and a loss of 11q were in agreement with our findings. Finally, 8p22-21 loss in erb2-amplified tumors was also seen by The Cancer Genome Atlas Network.<sup>28</sup>

Some studies have characterized ER-positive and ER-negative tumors by aCGH (Loo et al<sup>29</sup>) or by conventional CGH.<sup>30</sup> Although this is a more rough differentiation using only two subgroups, many of their results were in agreement with our findings. Loo et al<sup>29</sup> identified a gain of 3q25-27 and losses of 3p21, 4p16, 5q, 10q25, 14q32, 16p13, and 17p11 in ER-negative tumors, which is in agreement with our findings for basal-like tumors. Among the ER-negative tumors, they also identified a gain of 17q12 and losses of 8p21 and 17p11, supporting our findings for ERBB2-like tumors. For receptor-positive tumors, a gain of 16p11-13 and a loss of 16q13-24 was in agreement with our results for luminal tumors.

Altogether, many of the regions we have identified are supported by other studies measuring the DNA copy number by more direct techniques, as compared to our indirect expression-based method. This strongly supports our conclusions and also justifies using this approach for mechanisms within subgroups. We identified many regional aberrations differing between subgroups that have not been reported before. This may be explained by our different approach (including using gene expression data instead of directly measuring DNA copy numbers, for example, by aCGH). This has allowed us to considerably increase the sample size because only few small aCGH datasets with relevant information are available.

We have refined the data included in our analysis to obtain rates of distant metastasis or death from breast cancer as the outcome, but for one dataset (Hu), only time to relapse was available. However, this only introduces a minor bias because the majority of the typically observed relapse events included metastasis, and because only one of the six training datasets had this outcome definition. Another potential bias is the inclusion of patients that have received adjuvant treatment, which will bias the conclusion about metastasis biology because some patients will change their outcome status because of the treatment. However, the fraction of patients that respond to treatment and experience good outcomes as a consequence of treatment will typically be below 10% (adjuvant online, http://www.adjuvantonline.com), meaning that this will also be a minor bias. A varying fraction of the tumors in the individual datasets have disseminated cells to the lymph nodes. The classification of lymph node-positive patients without recurrence as nonmetastasis may be controversial. This may bias the results towards the metastatic mechanisms following primary spread to the lymph node. We accepted these biases in the study design to obtain a broad representation of tumor subtypes; the exclusion of lymph nodepositive and treated patients would reduce the number of ERBB2-positive and basal-like tumors markedly, resulting in a lack of power in these groups.

Combining the datasets by standardization, as we have done for the survival analysis, assumes that there are comparable characteristics in the included datasets. The varying fraction of node-positive tumors violates this assumption. However, this results in a more conservative test of gene sets in the survival analysis. This may explain why only four of the nine gene sets identified in the meta-analysis were significant in the survival analysis of the training set. Furthermore, these four gene sets are all significant in the test set, even though the sample size is considerably smaller. This is in agreement with the better separation of survival curves and the higher hazard ratios in the test set (Figs. 2C, G, 3C, and G). This may be explained by the very homogeneous test set, which included only node-negative samples from untreated patients performed with the same microarray platform.

## Conclusion

By performing a pathway meta-analysis, we have demonstrated that the previously described molecular subtypes account for the majority of metastatic mechanisms in breast cancer. However, by performing the analysis within these subgroups we have identified two different regions on chromosome 5q that impact metastasis among patients with basal-like and ERBB2-amplified tumors, respectively. We suggest one gene, *HAPLN1* at 5q14, which may potentially be a metastasis gene for basal-like tumors. The impact of these regions needs to be validated in future large studies using more direct measures of the DNA copy number (for example, aCGH). Furthermore, the role of *HAPLN1* and other potential driver genes needs to be addressed in further functional studies. We have also identified two pathways, TNFR2 and IL-12, which are involved in the metastasis of basal-like and ERBB2-amplified breast cancer, respectively. The molecular subgroups already form the basis for different treatments (such as trastuzumab, which is used against ERBB2-amplified tumors); however, future treatment protocols will aim to establish even more individualized strategies. Our study has identified genomic regions, single genes, and pathways that may be potential targets for future drug design for certain subgroups of patients.

## Methods

#### Data sets

Eight publicly available datasets examining gene expression at the ribonucleic acid (RNA) level in primary tumors were included in the analysis. These studies were performed with different platforms, different populations, and so on, as depicted in Table 1. The outcome used is distant metastasis or death from breast cancer, which is nearly always caused by distant metastasis. Only one data set (Hu) included local and regional recurrences. However, nonmetastatic relapse constitutes a minority of clinical cohorts. For the TRANSBIG dataset, samples from Sweden were removed to avoid sample overlap with the Uppsala and Stockholm datasets. The resulting dataset is termed TRANSBIG-S.

The normalizations performed in the studies were retained because the authors found these methods optimal for the datasets, and because the pathway analysis was performed separately in each dataset.

## Molecular subtypes

To identify the molecular subtypes, a single sample predictor was applied as described.<sup>8</sup> Prior to this, data were preprocessed within each dataset as follows. First, probe sets with maximal expression values were selected whenever more probe sets recognized the same gene using the "collapse to gene symbol" function in GSEA. Data were then column standardized for each sample by subtracting the mean expression of all genes in that sample from each genes expression value, and dividing by the standard devia-



tion for that sample. Next, row median centering was performed within each dataset by subtracting the median expression for a gene across samples from all expression values for that gene. Pearson's correlation coefficient between each sample and each of the five centroids (defined by Hu et al<sup>8</sup>) were calculated, and the sample was assigned the subtype with highest correlation coefficient. If the correlation coefficient was below 0.1 for any of the centroids, the sample was not assigned a subtype. Using this method, the samples were forced into the centroids defined by Hu et al.<sup>8</sup>

# GSEA analysis of pathways and genome regions associated with molecular subtypes

To analyze genome regions and pathways that were differentially expressed between the subtypes, we compared one subtype at a time with all other tumors. Only the seven datasets with successfully identified molecular subtypes were included in the analysis. For this analysis, we used original data (ie, not standardized). GSEA version 2.031 was used with 639 curated gene sets representing individual pathways. These pathway gene sets are adopted from KEGG (www.genome.ad.jp/KEGG), GenMapp (http:// www.genmapp.org), Biocarta (www.biocarta.com), and so on, and gathered in the Molecular Signature Database implemented in GSEA. Furthermore, we applied the analysis to positional gene sets delimited by cytobands downloaded from the Molecular Signature Database (http://www.broadinstitute.org/ gsea/msigdb/index.jsp).

The GSEA program ranks genes according to a signal-to-noise value:

$$(XA - XB)/(sA + sB), \tag{1}$$

where X is the mean and s is the standard deviation for the two classes A and B (one subtype and the remaining tumors, respectively). When several probes recognized the same gene, the probe with the maximum expression value was extracted using the "collapse to gene set" function. Gene sets represented by less than 15 genes in a dataset were excluded. The output from GSEA is an enrichment score, describing the imbalance in the distribution of ranks of gene expression in each gene set between the compared groups. The enrichment



score is normalized according to the size of the gene sets. Then, the gene sets were ranked according to the normalized enrichment score, with gene sets upregulated in the subgroup of interest on the top and downregulated gene sets at the bottom.

#### GSEA meta-analysis

The ranked lists of gene sets for each analysis generated by GSEA from the seven datasets were integrated so that only gene sets represented in the output from all datasets were included. The initial 639 pathway gene sets were reduced to 347 gene sets passing the threshold (at least 15 genes in gene sets) in all datasets. For the analysis of chromosomal regions, 386 chromosomal gene sets from the Molecular Signature Database were reduced to 188 gene sets.

For each dataset, individual gene sets were assigned a ranking value from 1 to the maximum number of gene sets, according to the ranking performed by GSEA. The mean ranking value for each gene set was calculated across the datasets and, finally, the gene sets were ranked according to this value.

Our null hypothesis is that the expressions of genes in the pathway gene sets are unrelated to the molecular subtype. This means that the ranking value for a given gene set in a given dataset is expected to be a random value between 1 and the maximum number of gene sets analyzed. To simulate the distribution of the mean ranking values across the six test datasets assuming the null hypothesis, a random drawing of the six ranking values was performed 10<sup>6</sup> times, and the mean value was calculated each time. A null distribution of the mean ranking values was generated from these results. To test the significance for a given gene set, the observed mean ranking value was compared to the null distribution. To fulfill the null hypothesis, an observed mean ranking value should be within 95% of the interval of the null distribution. Correction for multiple testing was performed by calculating the FDR, controlling the expected proportion of incorrectly rejected null hypotheses. These calculations were performed in the R environment (http:// www.bioconductor.org). Gene sets with FDR values below 0.01 were considered significant. Validation of the identified gene sets in the test set (Mainz data set) was performed by GSEA analysis using a nominal *P*-value

# Mechanisms associated with metastasis among each molecular subtype

To identify the chromosomal regions and pathways involved in metastasis within the molecular subtypes, GSEA analysis was performed as described above. Once again, genes were ranked according to the signal-to-noise value:

$$(XA - XB)/(sA + sB),$$
(2)

except that the compared groups were now tumors from patients that developed metastasis (class A) and tumors from patients that did not develop metastasis (class B). This analysis was performed within each subtype in each dataset, and the meta-analysis was applied to identify gene sets concordantly deregulated in metastasizing tumors across datasets for each molecular subtype. Again, 347 pathway gene sets and 188 positional gene sets passed the inclusion threshold. FDR values below 0.05 were considered significant.

#### Survival analysis

The above GSEA analyses were performed independently in each data set. To get a measure of the overall expression level for all genes in an identified region or pathway for each tumor—and in order to be able to compare these values across datasets-we calculated the region scores as follows. We started out with normalized data. First, probe sets with maximal expression values were selected whenever additional probe sets recognized the same gene using the "collapse to gene symbol" function in GSEA. Then, probe sets recognizing the same genes from the three different platforms were extracted using the gene symbol as an identifier, resulting in 6,654 annotated genes. Data were then row standardized within each dataset by subtracting the mean expression from each expression value for each gene, and dividing by the standard deviation for that gene. Finally, the standardized datasets were combined to a training set consisting of 1,239 tumors and a test set containing 347 tumors. The combination of row standardized datasets assumes that the sample characteristics (such as the clinical parameters) are comparable between the datasets. This introduced a bias in the training set because the dataset contained a variable fraction of lymph node-positive samples and patients receiving adjuvant treatment.

However, we retained the node-positive samples to obtain a reasonable number of ERBB2 and basal-like tumors. Patients with these tumors are more likely to have lymph node metastasis and are more likely to receive adjuvant medical treatment.

For the chromosomal regions and pathways, we calculated a score for each sample as the mean standardized expression values for all genes in the gene set. The score is thus a continuous variable representing the overall expression level in the region or pathway. The region score is related to the DNA copy number. However, using gene expression data alone, it is not possible to define a cut-off region in the score between different DNA copy numbers.

The association of region and pathway scores to survival was analyzed by Kaplan–Meyer plots and a log-rank test using surv and pchisq packages, respectively, in R. A high score (region score or pathway score) subset was defined as the third of samples having the highest score, and a low score subset included the third with the lowest score values. The middle third was excluded from the analysis. Hazard ratios were calculated by fitting a Cox model to the data, assuming a constant hazard ratio over time. The *P*-values were calculated using a log-rank test, with the null hypothesis postulating that there were no differences in survival between the groups.

# Validation of regions and pathways involved in metastasis

Validation of prognostic gene sets was performed in the latest published datasets (Mainz, TRANSBIG-S). The patients in these datasets are all node-negative and have not received adjuvant medical treatment; both studies were performed with Affymetrix HG133A chips minimizing bias from the treatment effect, clinical differences, and platform differences. For the validation of metastatic mechanisms in basal-like tumors. 41 basal-like tumors from the Mainz data were used. TRANSBIG-S data where the identification of molecular subtypes failed were excluded for that analysis. For ERBB2-amplified tumors, the number in the Mainz dataset was too small, and the TRANSBIG-S dataset was included, resulting in 347 samples. Instead of using single-sample prediction, we classified tumor ERBB2 status as 20% of tumors with the highest ERBB2 expression. The cut-off was set to



20% of tumors, because this resembles the frequency of ERBB2 amplification in breast cancer.

The region and pathway scores were calculated as described for the training set described above, and once again, the extreme groups (one-third of the highest scores and one-third of the lowest scores) were compared by survival analysis.

#### **Author Contributions**

Conceived and designed the experiments: MT, TAK. Analyzed the data: MT, QT. Wrote the first draft of the manuscript: MT. Contributed to the writing of the manuscript: MT, MB, QT, TAK. Agreed with manuscript results and conclusions: MT, MB, QT, TAK. Jointly developed the structure and arguments for the paper: MT, MB, QT, TAK. Made critical revisions and approved final version: MT, MB, QT, TAK. All authors reviewed and approved of the final manuscript.

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#### **Competing Interests**

Author(s) disclose no potential conflicts of interest.

## **Disclosures and Ethics**

As a requirement of publication the authors have provided signed confirmation of their compliance with ethical and legal obligations including but not limited to compliance with ICMJE authorship and competing interests guidelines, that the article is neither under consideration for publication nor published elsewhere, of their compliance with legal and ethical guidelines concerning human and animal research participants (if applicable), and that permission has been obtained for reproduction of any copyrighted material. This article was subject to blind, independent, expert peer review. The reviewers reported no competing interests.

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# **Supplementary Materials**

Direction in metastasis	Region or pathway	Mean ranking value	Max ranking value	P-value	fdr	Survival analysis <i>P</i> -value
Basal						
Up	5q14	26.8	188	0.0004	0.05	0.007
,	8g24	28.2	188	0.0005	0.05	0.17
Down	TNFR2	290	347	0.0002	0.04	0.009
ERBB2						
Up	16Q22	27.4	188	0.0004	0.05	0.15
	Xq22	28.4	188	0.0005	0.05	0.98
	Fructose mannose	32.5	347	0.00004	0.01	0.32
Down	5q33 —	165	188	0.0002	0.04	0.08
	IL-12	303	347	0.0002	0.04	0.004
	GPCRDB	313	347	0.00006	0.02	0.15

Table S1. GSEA meta-analysis within subtypes and survival analysis in training sets.





Figure S1. Distribution of ERBB2 and GRB7 expression in test set.

**Notes:** For validation of metastatic mechanisms in ERBB2 positive tumors, a test set including TRANSBIC-S and Mainz was used resulting in a total of 347 samples. Instead of using single sample prediction, we classified tumor ERBB2 status as 20% of tumors with highest ERBB2 expression.



Non-met	Met	Gene	P-va
		COX7C	0.
		TBCA	0.106
		AP3B1	0.388
		HOMER1	0.466
		SSBP2	0.16
		RPS23	0.021
		XRCC4	0.498
		HAPLN1	6.95E
		EDIL3	0.36
		MEF2C	0.191
		CETN3	0.935
		POLR3G	0.008
		CAST	0.798
		GLRX	0.611

Gene	P-value
COX7C	0.248
TBCA	0.106596
AP3B1	0.388029
HOMER1	0.466113
SSBP2	0.16987
RPS23	0.021021
XRCC4	0.498172
HAPLN1	6.95E-05
EDIL3	0.36699
MEF2C	0.191809
CETN3	0.935856
POLR3G	0.008516
CAST	0.798647
GLRX	0.611449

Figure S2. 5q14, 321 basale.