Distribution of the 21-Gene Breast Recurrence Score in Patients with Primary Breast Cancer in Germany

Verteilung des 21-Gen-Rezidiv-Scores bei Patientinnen mit primärem Mammakarzinom in Deutschland

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Key words
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Schlüsselwörter
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ABSTRACT

Background Multigene assays are being used increasingly to aid in decision-making about chemotherapy in breast cancer. Here, we present the 21-gene recurrence score (RS) of patients tested in routine clinical practice in Germany.

Patients and Methods In a retrospective analysis, 4695 patients with hormone receptor-positive and HER2-negative early breast cancer (pT1–3, pN0–1, M0) were included in whom RS testing was conducted in Germany between November 2015 and July 2018. RS groups as defined in the TAILORx trial (RS result 0–10; 11–25; 26–100) were used.

Results Of these patients, 21% were assigned to the low RS group, 63% to the midrange RS group, and 15% to the high RS group. 1772 (81%) of 2175 node-negative patients over 50 years of age were grouped either into the low RS group or the midrange RS group. The portion of patients with a low or midrange RS was 90% among node-positive patients (1284 of 1432 patients), 79% among patients with Ki-67-high (≥ 20%) tumors (1829 of 2310 patients), 86% vs. 70% among patients with G2 and G3 tumors (3244 of 3762 patients and 368 of 522 patients), respectively, 88% among patients with a tumor size of > 5 cm (140 of 159 patients), and 82% among node-negative patients at high clinical risk (1110 of 1352).

Conclusions The distribution of the 21-gene RS in German patients that were tested in routine clinical practice indicates that, according to the results of the TAILORx trial, chemotherapy may not be beneficial in most of these.

ZUSAMMENFASSUNG

Einleitung Multigen-Assays werden zunehmend als Entscheidungshilfe für eine Chemotherapie beim Mammakarzinom verwendet. Wir stellen hier den 21-Gen-Recurrence-Score (RS) von Patientinnen mit Brustkrebs vor, die routinemäßig in Deutschland untersucht wurden.


Ergebnisse Von diesen Patientinnen wurden 21% in die niedrige RS-Gruppe, 63% in die mittlere RS-Gruppe, und 15% in die hohe RS-Gruppe eingeteilt. 1772 (81%) von 2175 Patientinnen im Alter von über 50 Jahren und ohne Lymphknotenbefall wurden entweder in die niedrige oder die mittlere RS-
Gruppe eingeteilt. Der Prozentsatz an Patientinnen mit einem niedrigen oder mittleren RS betrug 90% bei Patientinnen ohne Lymphknotenbefall (1284 von 1432 Patientinnen), 79% bei Patientinnen mit einem hohen (≥20) Ki-67-Wert (1829 von 2310 Patientinnen), 86% bzw. 70% bei Patientinnen mit G2- bzw. G3-Tumoren (3244 von 3762 Patientinnen bzw. 368 von 522 Patientinnen), 88% bei Patientinnen mit einem Tumor durchmesser von >5 cm (140 von 159 Patientinnen), und 82% bei Patientinnen ohne Lymphknotenbefall, aber mit einem hohen klinischen Risiko (1110 von 1352).

**Ergebnisse** Die Verteilung des 21-Gens RS bei deutschen Patientinnen, die in der klinischen Routinepraxis getestet wurden, deutet darauf hin, dass gemäß den Ergebnissen der TAILORx-Studie die Chemotherapie bei den meisten dieser Patientinnen keinen Nutzen hat.

**Abbreviations**

HER2  human epidermal growth factor 2  
HR  hormone receptor  
IQR  interquartile range  
RS  21-gene recurrence score

**Key Message**

The 21-gene breast recurrence score classified 83% of node-negative and 90% of node-positive patients tested in routine clinical practice in Germany as low or midrange RS.

**Introduction**

Breast cancer is the most common cancer and remains the number one cause of cancer-related deaths among women in the US [1] and Europe [2]. With advances in diagnostics and therapy, however, breast cancer mortality has improved remarkably over the last few decades [1]. Amid these developments, increasing efforts are being made to distinguish between patients who are likely to benefit from adjuvant chemotherapy and those who can be spared the toxic side effects while retaining their favorable prognosis [3, 4].

As weighing the advantages and disadvantages of chemotherapy is challenging, especially in hormone receptor (HR)-positive, human epidermal growth factor 2 (HER2)-negative patients [5], a number of multigene assays, such as the 21-gene Oncotype DX breast recurrence score (Oncotype DX®, RS) [6], the 70-gene signature (MammaPrint®) [7], Endopredict® [8], and Prosigna® [9] are used in routine clinical practice to aid in decision-making.

The TAILORx trial was designed to prospectively validate the ability of the RS to estimate chemotherapy benefit in axillary lymph node negative HR+ HER2− patients [10]. Here, patients with a low RS (≤10) were assigned to receive endocrine treatment alone and patients with a high RS (≥26) were assigned to receive chemoendocrine treatment [10]. Patients with a midrange RS between 11–25 were randomized to receive either chemoendocrine treatment or endocrine treatment alone [10]. Initially published results show 9-year distant recurrence risks of 5, 8, 7 and 15% for the low RS, midrange RS + endocrine therapy, midrange RS + chemoendocrine therapy and high RS groups, respectively, and non-inferiority of outcome in the midrange RS group not receiving chemotherapy on the basis of the RS result was postulated with some exceptions [11].

These exceptions were described in more detail in a secondary analysis recently published, where it was shown that clinical risk as defined in the MINDACT (Microarray in Node-Negative Disease May Avoid Chemotherapy) trial [12] (low clinical risk if primary tumor ≤3 cm & low grade or ≤2 cm and intermediate grade or ≤1 cm and high grade) provided additional prognostic information in all RS groups [13]. Furthermore, patients ≤50 years of age seemed to benefit from chemotherapy if their RS was 21–25 or if they were at high clinical risk with an RS of 16–20 [13].

Retrospective analyses of several prospective trials have suggested that the RS is prognostic and predictive of chemotherapy benefit also in node-positive patients [14–16]. According to the 2018 German S3 breast cancer guidelines, as well as the 2019 AGO breast cancer guidelines, multigene assays may be used for patients with HR-positive HER2-negative, node-negative disease (in case of Oncotype DX, Prosigna and Endopredict) or in N0–N1 patients irrespective of hormone receptor and HER2 status (in case of MammaPrint), only if no clear decision regarding the use of adjuvant chemotherapy can be made based on conventional prognostic parameters [17, 18]. The current NCCN guidelines strongly recommend considering the 21-gene assay in HR-positive node-negative patients with tumors >0.5 cm in size and to consider multigene assays in HR-positive node-positive patients with <4 involved lymph nodes [19].

Here, we compare the RS result with clinical parameters in patients in Germany with primary invasive breast cancer for whom Oncotype DX testing was performed in routine clinical practice to aid in treatment decision-making.

**Methods**

**Patients and recurrence score**

This is a retrospective analysis of patients with HR-positive and HER2-negative early invasive breast cancer (pT1–3, pN0–1, M0) who received an Oncotype DX test in routine clinical practice between October 2015 and June 2017 in Germany. For this purpose, we obtained retrospective, anonymized data from Genomic Health Inc., Redwood City, USA. Grading and Ki-67 were evaluated by local pathologists and submitted to Genomic Health Inc. alongside the patients’ lymph node status, tumor size and age. We could not obtain data about the treating entity or therapeutic regimen. After being comprehensively informed by their treating physicians, patients who underwent RS testing had to sign an informed Consent Document providing detailed explanations about the purposes and use of personal data comprising

**Introduction**

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scientific research and related publications. The Informed Consent Document is regularly updated and complies with all applicable data protection laws, regulations and rules; in particular the EU-GDPR and the German Federal Data Protection Act. No ethics vote was required for analyzing the anonymized data according to the ethics commission of the University Hospital Tübingen, Germany.

Statistical analysis

We defined the low, midrange, and high RS groups in accordance with the definitions used for the TAILORx trial as an RS of 0–10, 11–25, and 26–100, respectively. Additionally, Ki-67 values of <20% were defined as low and ≥20% as high [20]. For clinical risk, the definition used in the MINDACT trial [12] (low clinical risk if node-negative & primary tumor ≤ 3 cm & low grade or node-positive & ≤ 2 cm & low grade or node-negative & ≤ 2 cm and intermediate grade or node-negative & ≤ 1 cm and high grade) was deployed.

Correlations between categorical variables were assessed using Pearson’s Chi-Squared Test. The significance level was set at α < 0.05. All tests were carried out as two-sided. Statistical analysis was performed using R version 3.5.0 and data visualization using the ggplot2 package version 3.1.0.

Results

Patient characteristics

Data from 4695 HR-positive, HER2-negative patients were available for analysis. The mean age was 56.6 years (standard deviation 10.4 years) with a median tumor size of 1.8 cm (interquartile range [IQR] 1.3–2.5 cm) and a median Ki-67 of 20% (IQR 10–25%), placing 2177 (49%) into the low Ki-67 and 2310 (51%) into the high Ki-67 group. In all, 3263 (69%) patients had no lymph node involvement, while 1432 (31%) were node positive. Furthermore, 283 (6%), 3762 (82%), and 522 (11%) patients were graded as G1, G2, and G3, respectively and 1792 (40%) were classified as low clinical risk. Compared with node-negative patients, the node-positive patients had a higher proportion of lower grade, Ki-67-low, clinically high-risk, and larger tumors, all p < 0.001 (▶ Table 1).

Recurrence score

The RS distribution can be seen in ▶ Fig. 1. The median RS was 16 (IQR 11–22), placing 1003 (21%) patients into the low RS group, 2975 (63%) into the midrange RS group, and 717 (15%) into the high RS group. The distribution of clinicopathological patient characteristics by RS group is illustrated in ▶ Table 2. Patients with high-grade tumors, Ki-67-high tumors, and node-negative patients were more likely to be in the high RS group (all p < 0.001) whereas no association was seen for tumor size (p = 0.265) and clinical risk (p = 0.255) (▶ Table 2, ▶ Fig. 2). This was true both for patients older than 50 years and patients 50 years of age or younger. In the subgroups of node-negative and node-positive patients the same associations between clinicopathological features and RS result were observed as in the combined cohort (▶ Fig. 3). In 55% of cases, Ki-67 and RS concordantly classified patients as low – low/midrange or high – high (▶ Fig. 4); 1772 (81%) of 2175 node-negative patients over 50 years of age were assigned a low or midrange RS. The proportion of patients with a low/midrange

| ▶ Table 1 Patient characteristics by axillary lymph node status. |
|------------------|------------------|------------------|------------------|------------------|
|                  | All patients     | Axillary lymph node | p (χ2)          |
|                  | n               | negative n         | positive n      |
| All patients     | 4695 (100)      | 3263 (100)         | 1432 (100)      |
| Grading          |                 |                   |                 |
| G1 (%)           | 283 (6)         | 142 (5)            | 141 (10)        | <0.001           |
| G2 (%)           | 3762 (82)       | 2558 (81)          | 1204 (84)       |
| G3 (%)           | 522 (11)        | 440 (14)           | 82 (6)          |
| Ki-67            |                 |                   |                 |
| Low (%)          | 2177 (49)       | 1274 (41)          | 903 (64)        | <0.001           |
| High (%)         | 2310 (51)       | 1812 (59)          | 498 (36)        |
| Tumor size       |                 |                   |                 |
| ≤ 2 cm (%)       | 2644 (59)       | 1912 (62)          | 726 (51)        | <0.001           |
| 2–5 cm (%)       | 1703 (38)       | 1076 (35)          | 627 (44)        |
| > 5 cm (%)       | 159 (4)         | 94 (3)             | 65 (5)          |
| Age (years)      |                 |                   |                 |
| ≤ 50 (%)         | 1497 (32)       | 1087 (33)          | 410 (29)        | 0.002            |
| > 50 (%)         | 3197 (68)       | 2175 (67)          | 1022 (71)       |
| RS group         |                 |                   |                 |
| Low (%)          | 1003 (21)       | 641 (20)           | 362 (25)        | <0.001           |
| Midrange (%)     | 2975 (63)       | 2053 (63)          | 922 (64)        |
| High (%)         | 717 (15)        | 569 (17)           | 148 (10)        |
| Clinical risk    |                 |                   |                 |
| Low (%)          | 1792 (40)       | 1712 (56)          | 80 (6)          | <0.001           |
| High (%)         | 2686 (60)       | 1352 (44)          | 1334 (94)       |
### Table 2 Patient characteristics by RS group.

<table>
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<tr>
<th></th>
<th>All patients</th>
<th>low n (%)</th>
<th>midrange n (%)</th>
<th>high n (%)</th>
<th>p (χ²)</th>
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</thead>
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<td>717 (15)</td>
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<td>2053 (63)</td>
<td>569 (17)</td>
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<tr>
<td>positive</td>
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<td>362 (25)</td>
<td>922 (64)</td>
<td>148 (10)</td>
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<tr>
<td>Grading</td>
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<td></td>
<td></td>
<td></td>
<td>&lt;0.001</td>
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<tr>
<td>G1</td>
<td>283</td>
<td>86 (30)</td>
<td>181 (64)</td>
<td>16 (6)</td>
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<tr>
<td>G2</td>
<td>3762</td>
<td>805 (21)</td>
<td>2439 (65)</td>
<td>518 (14)</td>
<td></td>
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<td>G3</td>
<td>522</td>
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<td>285 (55)</td>
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<td></td>
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<td>1413 (65)</td>
<td>201 (9)</td>
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<tr>
<td>high</td>
<td>2310</td>
<td>394 (17)</td>
<td>1435 (62)</td>
<td>481 (21)</td>
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<td>&lt;2 cm</td>
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<td>1674 (63)</td>
<td>421 (16)</td>
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<tr>
<td>2–5 cm</td>
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<td>372 (22)</td>
<td>1095 (64)</td>
<td>236 (14)</td>
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</tr>
<tr>
<td>&gt;5 cm</td>
<td>159</td>
<td>38 (24)</td>
<td>102 (64)</td>
<td>19 (12)</td>
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<tr>
<td>Age (years)</td>
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<td></td>
<td></td>
<td>&lt;0.001</td>
</tr>
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<td>&gt; 50</td>
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<td>728 (23)</td>
<td>1949 (61)</td>
<td>520 (16)</td>
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<td>1093 (64)</td>
<td>281 (16)</td>
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<td>854 (63)</td>
<td>242 (18)</td>
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<td>1334</td>
<td>329 (25)</td>
<td>863 (65)</td>
<td>142 (11)</td>
<td></td>
</tr>
</tbody>
</table>

> Fig. 1 Distribution of RS in clinical routine in Germany.
RS was 90% among node-positive patients (1284 of 1432 patients), 79% among patients with Ki-67-high tumors (1829 of 2310 patients), 86% vs. 70% among patients with G2 and G3 tumors (3244 of 3762 patients and 368 of 522 patients), respectively, 88% among patients with a tumor size of > 5 cm (140 of 159 patients) and 82% of node-negative patients at high clinical risk (1110 of 1352).

Discussion

Using the RS in a large cohort of patients with HR-positive/HER2-negative early breast cancer in Germany, we could identify according to the RS a large proportion of patients with clinically high-risk features such as high Ki-67 or high tumor grade, who according to the TAILORx results do not benefit from additional chemotherapy [11, 13]. This benefit may likely be attributed to ovarian suppression caused by chemotherapy treatment [13] and should be investigated in future trials. Considering "prospective retrospective" data [21] from the National Surgical Adjuvant Breast and Bowel Project (NSABP) B20 [22] for chemotherapy benefit in patients with a low RS and TAILORx data [11,13] for midrange RS patients, women > 50 years of age with an RS ≤ 25 and women ≤ 50 years of age with an RS ≤ 15 regardless of clinical risk or ≤ 20 at low clinical risk do not seem to benefit from chemotherapy in addition to endocrine therapy. However, subgroup analyses revealed that women ≤ 50 years of age with an RS of 21–25 regardless of clinical risk and women ≤ 50 with an RS of 16–20 at high clinical risk may in fact derive a benefit from additional chemotherapy [13]. In the prospective Plan B study, clinically high-risk patients (including patients with 1–3 involved lymph nodes) with low RS (≤ 11) had an excellent prognosis (94% 5-year DFS), although chemotherapy benefit was not evaluated separately within the node-positive patients. Retrospective data from prospective trials have been published indicating that node-positive patients with a low recurrence score may also not benefit from chemotherapy [14–16]. In the prospective Plan B study, clinically high-risk patients (including patients with 1–3 involved lymph nodes) with low RS (≤ 11) had an excellent prognosis (94% 5-year DFS), although chemotherapy benefit was not evaluated separately within the node-positive patients.
Additionally, the MINDACT trial has prospectively shown that multigene assays can be of help in guiding therapy decisions regardless of node status [12]. There, 1,404 node-positive patients were included and 912 (65%) were classified as low genomic risk [12]. Of the 709 node-positive patients at high clinical and low genomic risk, 356 received no chemotherapy and no statistically significant difference was seen in distant metastasis free survival when compared with the 353 patients, who received chemotherapy [12]. Results from the RxPONDER study (NCT01272037), which aims to determine the clinical validity of OncotypeDX in node-positive patients, are currently pending. This prospective randomized trial is set to define cut-off values for the possible omission of chemotherapy in node-positive patients [24]. As these are not yet available, we used the TAILORx cutoff values of 0–10, 11–25, and 26–100 for our entire cohort, which includes node-positive patients of whom 90% had an RS $\leq$ 25.

Ki-67 is used clinically to distinguish between luminal A- and B-like subtypes [20]. However, no clear cut-off values have been established; there is also high inter-observer variability and its role in predicting chemotherapy benefit remains unclear [25–27]. Using 20% as cut-off value, 49% of the patients in our cohort would have been classified as luminal A-like; however, 10% of these patients had a high RS, which is in line with earlier results [23]. Although we and others [28] found an association between Ki-67 and RS, the overall concordance rate of patients who would have received chemotherapy or not according to Ki-67 and RS results, respectively, was 55%. What measure to base treatment decisions on, in such cases, where the RS and Ki-67 lead to different conclusions, is a question that has yet to be answered.

In retrospective analyses the RS has been shown to frequently disagree with other molecular tests [29] and in postmenopausal women in the TransATAC trial it was outperformed by other multigene assays as a prognostic tool, even after improving its performance substantially incorporating clinicopathological information in form of the RS-pathology-clinical assessment of distant recurrence risk (RSPC) [30]. However, the only two tests validated to estimate chemotherapy benefit (or rather the lack thereof) in large prospective trials are the 70-gene signature (MammaPrint) [12] and the RS [11, 13].

In the MINDACT trial, patients at low clinical but high genomic risk did not benefit from chemotherapy and patients at high clinical but low genomic risk may have [12]. With the RS, the predictive value in the low and high RS group remains unclear, as these patients were not randomly assigned to treatment groups in the TAILORx trial but treated uniformly [10]. Further research in this area is needed and it therefore remains important even in the age of multigene assays to always take clinical risk into consideration when decisions on treatment are made [31].

![Fig. 3 Distribution of RS groups by patient characteristics in node-negative patients (a) and node-positive patients (b).](image-url)
As a limitation of this study, the distribution of tumor characteristics is biased by the decision to use the RS and is therefore not representative for the whole population of HR-positive/HER2-negative patients with early breast cancer. This is most likely why patients in whom axillary lymph nodes were involved more frequently had low-grade tumors and therefore had a higher rate of a low/midrange RS than node-negative patients. Just as observed in the Surveillance, Epidemiology, and End Results (SEER) database, node-positive patients in whom the RS was ordered tended to otherwise have low-risk clinical features when compared to their node-negative counterparts [32]. Bello et al. recently found that the RS distribution does not differ between node-negative and -positive tumors [33]. Additional limitations include the retrospective design of our study, the lack of treatment information and follow-up data, as well as the fact that no additional clinical risk factors such as menopausal status or progesterone receptor status were provided. We can therefore neither report patient outcome nor treatment efficacy. Furthermore, the records we received were incomplete with missing tumor size in 189 cases, Ki-
67 in 208 cases, grading in 128 cases, and age in 1 case, which is why the clinical risk of 217 patients could not be classified.

**Conclusion**

In conclusion, the RS was performed in routine practice in both node-negative and node-positive patients. In a large fraction of node-negative patients it indicates that chemotherapy may not be beneficial based on TAILORx results [11, 13]. In node-positive patients, its use was increased when other clinical factors, such as grading, Ki-67, or tumor size, indicated a lower clinical risk. Data from the prospective randomized RxPONDER-trial are awaited to evaluate whether patients with a low RS also might not benefit from chemotherapy and to determine the optimal RS cut-off values for decision-making.

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**Conflict of Interest**

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A. D. H. received a research grant, speaker and consultancy honoraria from Genomic Health Inc. and speaker fees and honoraria from Pfizer, Roche, Novartis, Lilly, MSD and AstraZeneca, Tesaro, Colovis and Eisai.

All remaining authors declare that they have no conflict of interest.

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