Comprehensive evaluation of methods to assess overall and cell-specific immune infiltrates in breast cancer

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Abstract

Background: Breast cancer (BC) immune infiltrates play a critical role in tumor progression and response to treatment. Besides stromal tumor infiltrating lymphocytes (sTILs) which have recently reached level 1B evidence as a prognostic marker in triple negative BC, a plethora of methods to assess immune infiltration exists, and it is unclear how these compare to each other and if they can be used interchangeably.

Methods: Two experienced pathologists scored sTIL, intra-tumoral TIL (itTIL), and 6 immune cell types (CD3+, CD4+, CD8+, CD20+, CD68+, FOXP3+) in the International Cancer Genomics Consortium breast cancer cohort using hematoxylin and eosin-stained (n = 243) and immunohistochemistry-stained tissue microarrays (n = 254) and whole slides (n = 82). The same traits were evaluated using transcriptomic- and methylomic-based deconvolution methods or signatures.

Results: The concordance correlation coefficient (CCC) between pathologists for sTIL was very good (0.84) and for cell-specific immune infiltrates slightly lower (0.63–0.66). Comparison between tissue microarray and whole slide pathology scores revealed systematically higher values in whole slides (ratio 2.60–5.98). The Spearman correlations between microscopic sTIL and transcriptomic- or methylomic-based assessment of immune infiltrates were highly variable (r = 0.01–0.56). Similar observations were made for cell type-specific quantifications (r = 0.001–0.54). We observed a strong inter-method variability between the omics-derived estimations, which is further cell type dependent. Finally, we demonstrated that most methods more accurately identify highly infiltrated (sTIL ≥ 60%; area under the curve, AUC, 0.64–0.99) as compared to lowly infiltrated tumors (sTIL ≤ 10%; AUC 0.52–0.82).

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Conclusions: There is a lower inter-pathologist concordance for cell-specific quantification as compared to overall infiltration quantification. Microscopic assessments are underestimated when considering small cores (tissue microarray) instead of whole slides. Results further highlight considerable differences between the microscopic-, transcriptomic-, and methylomic-based methods in the assessment of overall and cell-specific immune infiltration in BC. We therefore call for extreme caution when assessing immune infiltrates using current methods and emphasize the need for standardized immune characterization beyond TIL.

Keywords: Immune infiltrate, Breast cancer, Benchmarking, Methodology, Tumor infiltrating lymphocytes, Transcriptome, Methylome, Microscopy, Digital pathology

Background

In breast cancer (BC), the presence of immune infiltrate and its composition affects prognosis and treatment efficacy, including response to novel immunotherapies [1–5]. Specifically, increased levels of stromal tumor infiltrating lymphocytes (sTILs) are associated with response to neoadjuvant chemotherapy and prognosis in triple negative BC (TNBC) patients [4, 6–11]. In this context, sTIL has now been recognized as a valid prognostic biomarker by the expert panel of the 16th St. Gallen Breast Cancer Conference. Clinical trials investigating immunotherapies in BC are also using TIL or CD8+ T cell scores either for screening patients (e.g., NCT02997995) or as an endpoint (e.g., NCT03875573, NCT03815890, NCT03395899). Reliable methods to estimate the amount and composition of the immune infiltrate are therefore critical, for cross-study comparisons and future biomarker development.

Over the past years, several waves of technology have advanced the quantification and characterization of the immune infiltrate in solid tumors. Pathologists have developed methods to study the immune composition through the microscope [12, 13], while advances in computational biology have enabled the inference of cell type composition of solid tumors by utilizing bulk transcriptomic and methylomic data [14]. Currently, these methods are often used interchangeably in translational and fundamental research, assuming that they are providing similar information. However, the different methods do have different properties and it may be challenging to directly compare methods because of this. To the best of our knowledge, a detailed comprehensive comparison of these methods, including cell type inference using pathology, is still missing in BC, partly due to the lack of centralized microscopic, transcriptomic, and methylomic data.

The primary objective of the current study was therefore to compare the estimations of overall and cell-specific immune infiltration obtained by microscopic, transcriptomic, and methylomic methods in the International Cancer Genomics Consortium (ICGC) BC cohort [15–17]. The secondary objective was to evaluate the reliability of the different methods to classify tumors as highly or poorly infiltrated [10], as stratification of patients according to severity of sTIL infiltration has proven prognostic importance and may become a clinical biomarker in the near future [4, 10, 18].

Methods

Patients and dataset

This study is established on the ICGC BC cohort (https://dcc.icgc.org/) including 548 primary samples of female patients, for whom transcriptomic and methylomic data were available for 257 and 318 patients, respectively [15]. Data access was granted by ICGC. The generated pathology data, the type of data available for each patient, and the distribution of clinical and pathological characteristics are available at https://doi.org/https://doi.org/10.6084/m9.figshare.8234246.

Quantification of the tumor immune infiltrate

We collected tissue microarrays (TMAs; 3 cores/tumor, \( n = 254 \)), whole slides (WS) for IHC \(( n = 82, \) and hematoxylin and eosin-stained WS (H&E; \( n = 243 \)). TMAs and WS were stained for CD3, CD4, CD8, CD20, CD68, and FOXP3. Specific antibody clones, dilutions, and incubation times are listed in Additional file 1: Table S1. H&E slides were used to assess stromal TIL (sTIL) and intra-tumoral TIL (iTIL) for the whole slides and the TMAs. Two experienced pathologists (RS, HMH) scored all slides using the online pathology platform Slidescore [19] and reported the percentage of positive immune cells in the stromal and intra-tumoral compartment for each H&E- or IHC-stained slide, according to the existing guidelines [13]. A two-step digital image analysis (DIA) was performed using the Visiopharm Integrator System Software (VIS; Visiopharm A/S, Hoersholm, Denmark) using two optimized applications within the software to recognize positive DAB staining and tissue versus non-tissue. First, an application was used to detect the tissue and remove artifacts. With the second application, the positive-stained area (IHC-stained) was detected and the output variable is the positive area. Detection of, for example, CD3 positivity is based on the HDAB-DAB color deconvolution.
For all samples, the same threshold of positivity was kept.

For the computational analysis, we included methods that provided an estimation of the immune infiltration with respect to the entire tumor (Table 1, Additional file 1: Table S2). This list includes techniques used regularly for cell type inference. It should not be seen as exhaustive as new techniques are constantly being developed. We have included methods based on gene expression and methylation profiles, as both allow to study sample composition. Methods that estimate cellular populations from bulk data can be bluntly divided into two categories:

1. Those based on marker genes, providing an independent (semi-quantitative) assessment for each cell type, enabling comparison between samples but not within samples
2. Those based on deconvolution algorithms, inferring cell type fractions, enabling comparison between and within samples [14, 30]

To provide potential validation for immune infiltration, we calculated gene signatures specific for immune cell activity, namely cytolytic activity (CytAct) and interferon-gamma (IFNg) [31, 32]. All fractions of TIL and cell populations for each method are available at https://doi.org/https://doi.org/10.6084/m9.figshare.8234246.

Details for each method and processing are available in Additional file 1, which also includes Additional file 1: Tables S2 and S3.

### Statistical analysis

Statistical analyses were performed using R version 3.5.1. Values for immune infiltration were log transformed with an offset of 0.05. The agreement between observers and methods was assessed using the Bland-Altman method and Passing-Bablok regression analyses. Specifically, the geometric mean of the two scores from the same stained tumor section (x axis) is plotted against the ratio between the two methods or observers (y axis), considering the overall geometric mean of the ratios (center line) and the approximate 95% limits of agreement (horizontal lines), and Loess fitted curves were incorporated. Analysis was performed on all samples with available information for the methods taken into consideration. The concordance correlation coefficient (CCC) was used as a summary measure of reproducibility between observations [33] for each cell type. To establish the contribution of the iTIL and sTIL values to all TIL, a combined “averaged” TIL score was calculated by taking the arithmetic mean of the sTIL and iTIL scores for each sample and the Passing-Bablok regression was used for the comparison between TIL scores.

To assess replicability, correlations between methods for tumor immune infiltration were measured using the non-parametric Spearman’s rho coefficient, and a Loess smoothing was used for flexible interpolation. The 95% confidence intervals (CI) were calculated using the bootstrapping procedure with 1000 bootstrap samples (overlapping the results from the asymptotic approximation).

Both the Spearman’s rho coefficient and Lin’s CCC were interpreted according to qualifiers as “very poor”

### Table 1

| Method                              | Approach          | Description                                      | Overall immune score | Cell populations |
|-------------------------------------|-------------------|--------------------------------------------------|----------------------|------------------|
| H&E whole slide (WS)                | Microscopy based  | Pathology TIL scores                             | TIL scores           | NA               |
| IHC whole slide (WS)                | Microscopy based  | Pathology scores with IHC                        | Yes                  | NA               |
| Whole slide digital (digWS)         | Microscopy based  | Visiopharm digital scores IHC                    | NA                   | Yes              |
| H&E tissue microarray (TMA)         | Microscopy based  | Pathology TIL scores                             | TIL scores           | NA               |
| IHC tissue microarray (TMA)         | Microscopy based  | Pathology scores IHC                             | Summed lymphocyte fractions | Yes              |
| TMA digital (digTMA)                | Microscopy based  | Visiopharm digital scores IHC                    | Summed lymphocyte fractions | Yes              |
| Absolute CIBERSORT (aCBS) [20]      | Deconvolution     | Cell fractions, absolute mode used               | Summed lymphocyte fractions | Yes              |
| quanTIseq, lsft (qSEQ) [21]         | Deconvolution     | Cell fractions, absolute                        | Summed lymphocyte fractions | Yes              |
| MCP-counter (MCP) [22]              | Gene marker       | Arbitrary units                                  | NA                   | Yes              |
| xCell [23]                          | Gene marker       | Arbitrary units                                  | NA                   | Yes              |
| EPIC [24]                           | Deconvolution     | Cell fractions, absolute                        | Summed lymphocyte fractions | Yes              |
| MethylCIBERSORT (metCBS) [25]       | Deconvolution     | Cell fractions, absolute                        | Summed lymphocyte fractions | Yes              |
| TIL rna score (TILrna) [26]         | Gene marker       | TIL associated gene signature                    | TIL signature        | NA               |
| meTIL [27]                          | Gene marker       | TIL methylation profile                         | TIL signature        | NA               |
| Cell signatures Davoli et al. [28]  | Gene marker       | Computed gene signature                         | NA                   | Yes              |
| Cell signatures Danaher et al. [29] | Gene marker       | Computed gene signature                         | NA                   | Yes              |
(< 0.20), “poor” (0.20–0.40), “moderate” (0.40–0.60), “good” (0.60–0.80), and “very good” (0.80–1.00) [34].

To distinguish the highly and poorly infiltrated tumors, the (calculated) overall immune infiltration value (available at https://doi.org/10.6084/m9.figshare.8234246) from each method was utilized. The thresholds used for the categorization are based on the published meta-analysis on BC infiltration [10]: sTIL ≤ 10% for poorly infiltrated tumors, 11–59% for intermediately infiltrated tumors, and ≥ 60% for highly infiltrated tumors. To evaluate if different methods would classify the same tumor as poorly or highly infiltrated, receiver operator characteristic curves were generated and under the curves (AUCs) of each classifier were calculated using pROC package version 1.13.0. All tests were two-sided.

**Results**

**Assessment of immune cells on H&E and IHC whole slides**

Two experienced pathologists scored iTIL, sTIL, and six immune cell types (CD3+, CD4+, CD8+, CD20+, CD68+, FOXP3+) in an invasive primary BC cohort [15]. In line with previous reports [35–39], we observed a very good inter-observer CCC both for sTIL and iTIL (0.84 and 0.85, respectively, Fig. 1a). The limits of agreement showed a fair relative precision between measurements, and no major constant (intercept) or proportional (slope) drift between the two pathologists (Additional file 1: Figure S1). A very good concordance was observed between the stromal and averaged TIL (CCC 0.84, Fig. 1b), while by contrast, a poor concordance was observed between the intra-tumoral and averaged TIL score (CCC 0.37, Fig. 1c). These results show that immune infiltration in BC is mainly localized in the stromal compartment and is not greatly interfered by intra-
tumoral infiltration. Of note, some tumors with a high overall infiltration could still lack iTIL, as depicted in Fig. 1c, and include most probably tumors where the immune infiltrate is restricted to the stroma or to the tumor margin. The interobserver analysis demonstrated a moderate CCC for the immunohistochemical (IHC) assessments, where stromal scoring performed overall better and more precisely than intratumoral scoring (Fig. 1a). All stromal IHC CCC values were lower for immune cell subtypes than for sTIL and iTIL assessment on H&E, ranging from 0.63 to 0.66 (Fig. 1a). The intra-tumoral IHC methods had a CCC below 0.6, except for iTCD3 (CCC = 0.64). We therefore only considered the more reliable and abundant stromal estimates for further analyses.

Comparison of tumor immune infiltration estimates between tissue microarrays and whole slides (H&E and IHC)

When stromal TIL was scored on both whole H&E slides and TMA H&E, the correlation was moderate (0.56) (Fig. 3). The CCC between WS and TMA was only 0.26, where overall a lower (5.98× lower) percentage was scored on H&E TMAs compared to whole H&E slides (Additional file 1: Figure S2a). We characterized six immune features both on TMA and WS (Fig. 2a). The immune infiltrate scores were systemically higher on WS as compared to TMA, as depicted in Fig. 1d for the most abundant immune cell population, i.e., CD3+, and in Additional file 1: Figure S2a for the other immune cells. The CCCs between WS and TMA were globally poor and ranged between 0.21 for CD4+ and 0.43 for FOXP3+ cells (Additional file 1: Figure S2b). We further evaluated both TMA and WS using digital pathology and confirmed the higher level of immune infiltration estimated with WS as compared to TMA (Fig. 1e, Additional file 1: Figure S2c), highlighting the spatial heterogeneity of the tumor immune microenvironment (Additional file 1: Figure S2b). Figure 2b depicts one sample that shows high infiltration on the WS and limited infiltration in the TMA. In line with previous reports [35], we observed a moderate correlation between human and digital assessment of CD3+ (CCC = 0.42, Fig. 1f), yet the digital evaluation showed a lower estimation of all immune cells compared to pathologists.
suggesting a relative human overestimation or digital underestimation of tumor immune infiltration.

Comparison of microscopic, transcriptomic, and methylomic evaluation of overall tumor immune infiltration

To evaluate if different data types could estimate overall immune infiltration consistently, we compared the (calculated) score of several microscopic-, transcriptomic-, and methylomic-based methods. In addition, two inflammatory gene signatures (interferon-gamma and cytolytic activity) were calculated (pink category in Fig. 3), as these often correlate to tumor immune infiltration and can provide information on the status of the tumor microenvironment [31, 32]. A description of all methods is provided in Table 1 and Additional file 1: Table S2. First, while the Spearman correlations between microscopic (red) and all methylomic (blue) or transcriptomic (green) estimates were poor to moderate, our analysis showed that stromal infiltration correlates better with...
with all other methods, including transcriptomic and methylomic methods, as compared to the intra-tumoral infiltration (Fig. 3a). We tested the possibility that higher iTIL may lead to more pronounced inflammatory gene expression than sTIL as iTIL may have a crucial anti-tumor role [40–44]. Yet, we observed no higher correlation between intra-tumoral infiltration and inflammation-associated signatures (immune signatures, pink label Fig. 3a). Secondly, a good correlation was observed between stromal microscopy assessment on H&E-stained (sTIL) and IHC-stained WS, where the sum of T (CD3+) and B (CD20+) cells were considered ($r = 0.61$). Thirdly, the correlation with the immune gene signatures, cytolytic activity ($r = 0.51$) and IFNg ($r = 0.57$), improved slightly when the infiltrate was scored with digital pathology compared to the other microscopic (red) methods (Fig. 3a). This is most probably because both the immune signatures (CytAct and IFNg) and the digital assessment of the TMA do not consider the type of infiltrates (intra-tumoral vs stromal). The same trend was observed between the digital assessment of the TIL on TMA (digTMA) and transcriptomic TIL methods (green), where the Spearman correlations with EPIC ($r = 0.42$), aCBS ($r = 0.53$), and TILrna ($r = 0.56$) were again the highest compared to all other microscopic (red) methods. Fourthly, as expected, methods using the same modality for input of data showed better correlations. For example, several transcriptomic estimates showed a very good correlation with each other ($r > 0.80$), and methylCIBERSORT [25] and the methylomic TIL score, meTIL [27], showed a reassuring good agreement ($r = 0.77$). The correlation between transcriptomic and methylomics was variable, but methylCIBERSORT showed good correlations with absolute CIBERSORT ($r = 0.75$) and TILrna [26] ($r = 0.76$), and similar observations could be made for meTIL and CIBERSORT ($r = 0.66$) and meTIL and TILrna ($r = 0.70$). Fifthly, of the transcriptomic and methylomic methods, TILrna, methylCIBERSORT, and absolute CIBERSORT [20] showed the highest correlations with microscopy scores, though the correlations were still not perfect with the highest being 0.56 between digTMA and TILrna, 0.53 considering sTIL and methylCIBERSORT, and 0.53 between digTMA and absolute CIBERSORT. These numbers were however in line with a recently published lung cancer study [45]. Finally, of the methods that predict global immune infiltration based on the transcriptome (green label in Fig. 3a), TILrna showed the highest correlations with the various immune gene signatures (pink label, $r = 0.90–0.94$), while quanTIseq [21] showed the poorest correlations ($r = 0.16–0.18$). Similar analyses were further carried on separately for ER-negative and ER-positive tumors as the biological significance of the immune infiltrate may be different [46, 47] (Fig. 3b, c). The correlations for the microscopic-versus methylomic- and transcriptomic-based methods were in general slightly higher in the ER-negative compared to the ER-positive subgroup. Nevertheless, compared to all samples, the ER-negative tumors did not necessarily show higher correlations.

Following the analysis at the continuous level (Fig. 3a–c), we further aimed to investigate the ability of the methods to identify lowly infiltrated (sTIL $\leq 10\%$) and highly infiltrated (sTIL $\geq 60\%$) tumors as defined by Denkert and colleagues [10]. TIL is currently not used as a classifier in the clinic, yet the stratification of patients based on TIL has provided important prognostic information in clinical studies [4, 18]. To this end, a ROC curve analysis was performed (Fig. 4a, b) and most methods performed moderate to very poor in recognizing the lowly infiltrated tumors (Fig. 4a, c blue boxes). The iTIL score showed the highest area AUC for the lowly infiltrated tumors (Fig. 4c, blue boxes), and the methylomic-based TIL scores had a slightly higher AUC as compared to the transcriptomic-based methods, although their confidence intervals were overlapping. ROC analyses further showed that most methods can more accurately identify highly infiltrated tumors as compared to lowly infiltrated tumors (Fig. 4b, c red boxes). Here, the highest AUCs were still based on the microscopy methods, but methylomic- and transcriptomic-based methods also showed fair to high AUCs. Of interest, highly infiltrated tumors have the highest expression of inflammatory signatures, like interferon-gamma and cytolytic activity (Fig. 4d), yet lowly infiltrated tumors could also show a wide range of these inflammatory gene expressions.

**Comparison of microscopic, transcriptomic, and methylomic evaluation of cell-specific immune infiltration**

Several observations could be made from the comparison between microscopic and computational methods that quantify specific immune cell types with transcriptomic or methylomic data (Table 1, Fig. 5a–e). Firstly, the correlations between the microscopic evaluations and the transcriptomic and methylomic data were always $< 0.60$, with CD8$^+$ T cells showing the highest (Fig. 5a) and macrophages (Fig. 5e) the lowest correlation between microscopic and methylomic or transcriptomic methods, respectively. Of interest, there was no systematic increase in the correlation coefficients when considering WS instead of TMA scores. Of note, when we examined the CCC between methods for specific cell types, poor concordance (<0.3) was found for all methods (data not shown). Secondly, most omics-derived methods showed large inconsistencies between cell types regarding their correlation with microscopy. For example, while quanTIseq showed very poor correlation with sTIL (0.09, Fig. 3a) and CD4$^+$ cells (0.01, WS evaluation; Fig. 5b), the correlations for CD8$^+$ were
moderate (0.54, WS evaluation, Fig. 5a). Altogether, these analyses highlight the variable correlation between the different omics-derived methods and microscopic assessments, which further vary according to the immune cell type.

**Discussion**

In this study, we have comprehensively demonstrated that methods of the same modality (microscopy, transcriptomic, or methylomic based) to estimate overall infiltration show reassuring correlations, but the correlations deteriorate when comparing between modalities. In addition, we quantified specific immune cell types and observed a strong heterogeneity in the correlations between the microscopic- and omics-based estimates, and also between the different omics-based estimates. For each cell type, the best method to use may be different. Our analysis clearly shows that different transcriptomic and methylomic methods have limitations in estimating immune infiltration, as correlations with...
microscopy-based methods do not exceed 0.6. As the pathology assessment of TIL has reached level 1B evidence with new studies confirming its value for the patient [4], the fact that correlations are not high may warrant extra caution when using non-pathology methods as these may measure different characteristics of the immune infiltrate. These findings might be explained by the fact that transcriptomic and methylomic data are not perfectly representing protein expression of immune cells. In addition, regular bulk transcriptome or methylome analysis does not detect heterogeneity within a sample and ignores the localization of cells [48]. An example as depicted in Fig. 2b emphasizes the importance of spatial heterogeneity, but also raises the question about the functionality of the observed immune infiltrate. Importantly,
transcriptomic or methylomic methods may be biased towards a specific cell state, as they are often based on cells challenged with experimental processes (e.g., tissue digestion or flow cytometry) or from different origin (e.g., peripheral blood or different tumors), which may contribute to the observed discordance with microscopy.

More research into the contribution and spatial distribution of specific immune cells in the context of clinical outcome is warranted to understand beneficial and detrimental immune cell profiles. Spatially resolved methods, measuring genetic and phenotypic diversity, would support advances for clinical studies [49]. Moreover, integration of deep learning approaches with morphological features, in conjunction with genomic-derived data, will probably be needed to derive a full comprehensive evaluation of the immune environment in solid tumors. Our data provided us with insightful observations that can guide future research using digital image analysis. First, we showed that stromal evaluation of several immune cell types is reliable with an acceptable concordance observed between pathologists, especially for stromal infiltration. Second, values may also be estimated with a digital approach, taking into account the systematically lower numerical values for digital estimates [35]. Third, our results raise caution for the evaluation of immune markers on TMA (or biopsies) as small punches of the tumor may obscure information of infiltration and WS showed overall a higher infiltration than TMA. Our results on immune infiltrate are in contrast with several studies which demonstrated that TMA are reliable for the evaluation of several prognostic epithelial-based tumor markers [50–53].

Our second objective was to stratify patients into lowly infiltrated and highly infiltrated tumors, as more clinical studies showed that sTIL-based stratification could serve as an important prognosticator [4, 18]. We observed that microscopic methods (iTIL, WS, TMA IHC, digTMA) were better in the stratification of patients into lowly or highly infiltrates, as expected due to their common modality as sTIL (as opposed to, for example, gene expression-based methods). We also showed here that the majority of the methods are better in recognizing highly infiltrated tumors as compared to lowly infiltrated tumors. The lower accuracy for identification of lowly infiltrated tumors may be problematic as the majority of the breast tumors will have infiltration above 0% but far below 60%, as shown by Loi et al. [4] in early TNBC. These findings should be taken into consideration in developing inclusion and stratification criteria, as well as endpoints in the context of clinical trials.

Strengths of this study include the large number of patients and extensive central evaluation of immune cells, together with the availability of transcriptomic and methylomic data. A limitation of our study is that tissue analysis (FFPE) and DNA/RNA isolation (FF) were not performed on the exact same area from the tumor. Infiltration may be affected by heterogeneity and partially explain the correlations not exceeding 0.6. We minimized this effect by studying multiple cores spread throughout the tumor in the TMA and large sections of the tumors in WS, far exceeding the area usually evaluated with biopsies. Nevertheless, correlation with various omics-based methods did not systematically increase when considering WS versus TMA. In addition, the digital analyses of the immune cells were calculated as an area, while pathologists report a cellular percentage, leading to pre-analytical factors that affect the results. We were however not able to calculate the cellular percentage due to visual cell segmentation problems. Another limitation is that the omics-based methods considered in this manuscript do not consider the localization of the cells in the tumors. In this context, a recent study conducted in TNBC suggested that transcriptomics might have the potential to derive this spatial information [54]. Finally, this cohort does not provide follow-up or therapy response data. Future studies may test superiority of specific measures with the ultimate goal to better guide precision medicine.

**Conclusion**

This study highlights an important heterogeneity in the various estimates of immune infiltrates in BC and calls for caution when used in the clinical context. This study further provides an important resource of multi-level data of the tumor immune microenvironment to researchers for future investigations. Ultimately, there is an urgent need for the development of international guidelines to categorize breast tumors according to their immune infiltrate in both a quantitative and a qualitative manner. Combining the valuable information from multiple methods, e.g., the spatial information from pathology and transcriptomic information on cellular activity, may elucidate the role of immune infiltration in disease progression in a more accurate manner.

**Supplementary information**

Supplementary information accompanies this paper at https://doi.org/10.1186/s13058-019-1239-4.

**Additional file 1: Supplementary Methods. Figure S1.** The inter-observer analysis for all the stromal and intratumoral immune cell scores.

**Figure S2.** Comparison between (dig) WS and (dig)TMA. Table S1. The procedures, clones, manufacturer and dilution used for the immunohistochemistry on tissue micro array (TMA) and Whole Slides (WS). Table S2. Description of all methods used in the manuscript to estimate immune cell infiltration. Table S3. Overview of used cell fractions for overall immune infiltration and specific immune subtypes.

**Abbreviations**

AUC: Area under the curve; BC: Breast cancer; CCC: Concordance correlation coefficient; ER: Estrogen receptor; FF: Fresh frozen; FFPE: Formalin-fixed paraffin-embedded; H&E: Hematoxylin and eosin; HER2: Human epidermal growth factor receptor 2; ICGC: International Cancer Genomics Consortium; IHC: Immunohistochemistry; iTIL: Intra-tumoral tumor infiltrating lymphocyte; sTIL: Stromal tumor infiltrating lymphocyte; TIL: Tumor infiltrating
lymphocyte; TMA: Tissue microarray; TNBC: Triple negative breast cancer; Tregs: Regulatory T cells; WS: Whole slide

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Authors’ contributions

CD, EB, RS, HH, and IN conceived the study. GKH and KW selected the tumor blocks, and constructed and stained the TMAs. RS and HH did all the anatomic pathological scoring on the slides. JH provided an online platform to enable pathology scoring. LB and IN generated, interpreted, and supervised the digital analysis of the tissue slices. YB, BN, JK, and DB processed the transcriptomic and methylation data for the various methods. DB performed the statistical analysis under the supervision of EB and CD. DB, EB, CD, and IN interpreted the data. IN drafted the manuscript under the supervision of EB and CD. All other authors contributed data and/or materials; all authors critically commented and approved the final version.

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Availability of data and materials

The clinical, pathologic, transcriptomic, genomic, and methylation data used in this study was already publicly available upon request to the ICGC data access committee (https://icgc.org/daco/). The microscopic data that were generated specifically for this study are included in this published article and its additional information files.

Ethics approval and consent to participate

This ICGC Breast Cancer dataset is hosted by the ICGC and deposited at https://dcc.icgc.org/. Data access was granted by the International Cancer Genome Consortium (ICGC).

Consent for publication

Not applicable

Competing interests

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