Development of automated quantification methodologies of immunohistochemical markers to determine patterns of immune response in breast cancer: a retrospective cohort study

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ABSTRACT

Introduction: Lymph nodes are one of the main sites where an effective immune response develops. Normally, axillary nodes are the first place where breast cancer produces metastases. Several studies have demonstrated the importance of immune cells, especially dendritic cells, in the evolution of breast cancer. The goal of the project is to identify differences in the patterns of immune infiltrates, with particular emphasis on dendritic cells, in tumour and axillary node biopsies between patients with and without metastases in the axillary nodes at the time of diagnosis. It is expected that these differences will be able to explain differences in survival, relapse and clinicopathological variables between the two groups.

Methods and analysis: The study will involve 100 patients diagnosed with invasive breast cancer between 2000 and 2007, 50% of whom have metastases in the axillary lymph node at diagnosis. In selected patients, two cylinders from biopsies of representative areas of tumour and axillary nodes (with and without metastasis) will be selected and organised in tissue microarrays. Samples will be stained using immunohistochemical techniques for different markers of immune response and dendritic cells. Two images of each cylinder will be captured under standardised conditions for each marker. Each marker will be quantified automatically by digital image procedures using Image-Pro Plus and Image-J software. Associations of survival, relapse and other clinicopathological variables with the automatically quantified levels of immune infiltrates in patients with and without axillary node metastasis will be sought.

Ethics and dissemination: The present project has been approved by the Clinical Research Ethics Committee of the Hospital Universitari Joan XXIII (Ref: 22p/2011). Those patients whose biopsies and clinical data are to be used will give their signed informed consent. Results will be published in peer-reviewed journals.

INTRODUCTION

Breast cancer is the main cause of mortality from malignant tumours in women in western developed nations. In recent years, several strategies and therapies have been developed to cure patients or prolong their survival. However, some of these therapies have side effects and, in patients with distal metastasis, rarely have a curative effect. Several strategies have recently been used to develop treatments that are more effective and less harmful to patients. Therapeutic
Dendritic cell (DC) vaccines are being studied in several tumours and in breast cancer. The results obtained so far have not been completely satisfactory. DCs are the most important antigen-presenting cells (APCs). They originate in the bone marrow then move into the bloodstream and migrate to the peripheral tissues. At that point DCs are immature and carry out sentinel functions, waiting for signals—normally infection or damage that promotes inflammation—that trigger their maturation. During this period DCs are very effective at catching and processing protein antigens from the environment, but remain relatively inactive as APCs. At the appropriate time they migrate into the lymphatic vessels and travel to the regional lymph nodes, where they mature and present the antigens to T lymphocytes and activate the immune response. The main functions of DCs are: (1) to capture antigens and migrate to the lymphoid organs to optimise the clonal selection of CD4 and CD8 T cells; (2) to promote immune response-stimulating quiescent, and T and B memory lymphocytes; (3) to improve the T cell-mediated immune response; (4) to induce tolerance to the body’s own antigens.

Most DCs belong to a myeloid lineage and others to a lymphoid lineage. Those of the latter type are also known as plasmacytoid DCs. The various subpopulations of myeloid DCs (HLADR+CD11c+) can express markers such as CD1a (Langerhans cells), CD21 (follicular DCs), S-100 (interdigitating cells), while the plasmacytoid such as CD1a (Langerhans cells), CD21 (follicular lymphoid lineage. Those of the latter type are also immunologically competent and a potential site for the activation of specific T cells. On the other hand, the immune response dependent on DC-Lamp DCs is altered during metastasis formation, and is activated before and during the formation of micrometastases.

Nevertheless, DCs are involved in only one of the first steps in the antitumour immune response. If this step is altered, the subsequent immune response may not be appropriate. Thus, it is also necessary to evaluate what happens with the general immune response at the tumour and lymph node levels.

The recruitment of immune and/or inflammatory cells in invasive breast cancer is one of the procedures involved in the antitumour immune response. Studies have demonstrated the presence of a heterogeneous population of immune cells in the tumour microenvironment of the breast. Basically, this population comprises T lymphocytes (CD4, CD8), cytotoxic cells (T-cell intracellular antigen-1 (TIA)-1, granzyme B), tumour-associated macrophages (CD68), DCs (CD21, CD1a, CD123) and regulatory T cells (FOXP3). In some of these studies, the type and amount of these immune cells have been linked to clinical variables such as tumour size, age, disease progression and patient survival. The importance of these cellular infiltrates has also been used to predict clinicopathological performance. Moreover, the evaluation of components of the immune response from tumour and stroma could help differentiate the different types and stages of the illness. The presence of cells with immunosuppressive capacity in the tumour (regulatory cells), as well as
tolerance induction by tumour cells could explain why some tumours grow and metastasise progressively. A significant difference in the number of regulatory T cells has been detected between human epidermal growth factor receptor 2 (HER2)-positive and HER2-negative patients with advanced breast cancer.

In breast cancer, most research has partially addressed the subpopulations of immune cells present in the tumour or in the lymph nodes. Moreover, most of the cited articles have investigated only the tumour or the lymph nodes. Thus, it will be of interest to determine the global pattern of the immune response (DCs and other immune cells) simultaneously in breast cancer tumours and axillary lymph nodes to detect any differences associated with tumour evolution.

Most immune markers can be evaluated using immunohistochemistry (IHC), with which several cell subpopulations can be detected and quantified in situ in the tumour or lymph node. IHC is simpler, cheaper and faster than other molecular biological techniques. The markers can be evaluated quickly and efficiently through the use of tissue microarrays (TMAs), the standardisation of stain conditions of immunohistochemical markers, and the detection and quantification of markers by automated digital image analysis. Digital analysis can minimise the problems of human subjectivity between evaluators. In recent years, the use of digital image analysis of immunohistochemical markers in breast cancer has become important. Several systems, such as Automated Cellular Imaging System (ACIS) III (DAKO, Carpinteria), have been developed and commercialised and subsequently approved by the US Food & Drug Administration. Nevertheless, these systems only allow some types of immunohistochemical markers to be quantified, such as the membrane marker CerbB2, and nuclear markers, such as oestrogen and progesterone receptors.

The aims of this aspect of the proposed project are: (1) to develop new image analysis methods to quantify immune markers arising from other project activities; (2) to search for different patterns of immune response cells and specifically of DC infiltrates in the tumour and axillary lymph nodes, comparing patients with and without metastasis in the axillary lymph nodes; (3) to identify any associations between immune infiltrates and clinicopathological variables and patient survival.

METHODS AND ANALYSIS

Study design and setting
The study will examine a retrospective cohort of 100 patients with invasive breast cancer diagnosed in Hospital de Tortosa Verge de la Cinta, Catalonia, Spain, between 2000 and 2007. Fifty per cent of the patients had metastasis in the axillary lymph nodes at diagnosis. The amount and localisation of different types of immune infiltrates, especially of DCs, in tumour and in axillary lymph nodes will be evaluated in the patient biopsies. Immune infiltrates will be detected and evaluated using IHC and image analysis procedures. Differences in the patterns of immune infiltrates will be sought in the tumour biopsies and in axillary lymph nodes with and without metastases between the two groups of selected patients (with and without metastasis in axillary lymph nodes at diagnosis). Associations of survival, relapse and other clinicopathological variables with the immune infiltrate patterns will be sought.

Study period
The study will be undertaken between January 2012 and December 2014.

Study sample
The study will involve only the samples that fulfil the following inclusion criteria: invasive breast cancer samples with enough representative tumour (more than 50% of selected area must contain tumour) and axillary lymph node tissue in patient biopsies. Of these, only samples from patients who have given signed informed consent will be used.

Samples collected from 2000 to 2007 will be selected from the Tumour Bank of the Pathology Department of the Hospital de Tortosa Verge de la Cinta, as part of the XBTC included in the BioBanc node HTVC – Biobanc Hospital Universitari Joan 23 – IISPV. This period has been chosen to ensure enough samples with at least 5 years of follow-up in both patient groups studied (with and without axillary lymph node metastases).

Tumour and axillary node paraffin-embedded biopsies will be used to study the immune infiltrate in patients with breast cancer. In those without metastasis in axillary lymph nodes at diagnosis, two cylinders of 2 mm each from representative areas of the tumour and two from axillary lymph nodes will be taken. In patients with metastasis, six cylinders will be evaluated, two from the tumour, two from metastasis-free axillary nodes and two from axillary nodes with metastasis.

Sample size calculation
The study includes 100 patients, based on the sample size and power calculations for identifying relative risk available at http://www.imim.es/ofertadeserveis/software-public/granmo/. The calculation was performed to find an immune infiltrate pattern able to predict a death risk in the group with metastasis 1.6 times that of the group without metastases. Values used in the calculation are described below.

Accepting an α risk of 0.05 and a β risk of 0.2 in a two-sided test, 50 participants exposed to certain expected type of immune pattern and 50 non-exposed participants are necessary for a relative risk of death greater than or equal to 1.6 in the exposed group to be statistically significant. The proportion of deaths among the people in the non-exposed group has been estimated to be 0.5, based on the principle of maximum indeterminacy, since there are no previous data about the type of pattern of immune infiltrate that can predict death. A drop-out rate of 10% has been anticipated.
Samples and data processing

1. TMA construction: Twenty TMAs of 25 cylinders extracted from biopsies will be constructed. Eight TMAs will be constructed from tumour biopsies ((2 cylinders/patient)×100 patients)/(25 cylinders/TMAs). Eight TMAs will be obtained from axillary lymph nodes free of metastasis, and four will be from axillary lymph nodes with metastasis since only 50% of patients belong to this group.

2. Slide preparation: Eleven immune and DC markers will be evaluated by IHC. For this purpose, 11 sections of 2 μm thickness will be cut from each TMA to stain all markers. A total of 220 slides will be stained (20 TMAs×11 markers).

3. Clinical immune markers studied
   A. Markers for general immune response detection: CD4 (usually expressed in T helper lymphocytes), CD8 (usually expressed in cytotoxic T lymphocytes), CD57 (expressed in Natural Killer lymphocytes), FOXP3 (expressed in regulatory T cells), CD21 (expressed in mature B lymphocytes), CD68 (expressed on the surface of macrophages).
   B. Markers for DC detection: CD1a (dendritic Langerhans cells), CD123 (plasmacytoid DCS), S100 (interdigitating DCS) CD208 (DC Lamp+), CD68 (marker of mature DCs).

4. IHC staining of samples: slides will be stained fully automatically with the Autostainer Link 48 (DAKO) from the pathology department. The use of TMAs allows automation and the staining of a large number of samples quickly and under the same conditions. Quality controls for breast cancer used in the pathology department for daily clinical practice will be stained at the same time as the samples of the study. In this way, the positive and negative controls as well as the quality of the counterstaining may be compared with the IHC staining of the study samples. Moreover, all the TMAs will be stained for the same markers in the same time in order to ensure the same conditions of staining (same antibody dilution, same temperature, etc.).

5. Standardisation of image capture: To obtain maximum reproducibility in image analysis the conditions of image capture will be standardised for each marker, which means that we will have the same illumination conditions and a standard white balance.

   Two images from the area with the most abundant cellular infiltration of each cylinder will be captured, so by the end of this step we will have 11 000 digital images: ((200 tumoral cylinders×200 axillary lymph node without metastasis cylinder+100 axillary lymph node with metastasis cylinder))×(2 pictures)×(11 markers). As seen in our previous studies, two images taken from a tumour area have been demonstrative enough for immune cell infiltrate evaluation. Tissue sections will be viewed using brightfield illumination under a Leica DM LB2 upright light microscope (Leica Microsystems Wetzlar GmbH, Wetzlar, Germany) with a 40× plane-apochromatic objective with a numerical aperture of 0.63. The representative areas of the different sections will be captured on a Leica DFC320 digital camera (Leica Microsystems Digital Imaging, Cambridge, UK) controlled by Leica DFC Twain software V6.3.0 (Leica Microsystems Digital Imaging) running on a Compaq Professional Workstation computer (3.6 GHz Pentium IV CPU, 3.00 GB RAM). These images will have a resolution of 2088×1550 pixels with RGB 24 true colour format, and will be saved in uncompressed tagged-image file format (TIFF).

6. Development of image analysis procedures: Image analysis procedures for quantifying the 11 immunohistochemical markers will be adapted for each marker from our automated image analysis procedures developed in previous studies (see point 7). These methods will be validated by comparing the results obtained from several manual (pathologist) evaluations. Several strategies and image analysis techniques will be used:
   - Light image correction.
   - Determination of the most suitable space colour model (RGB, HSI, CMYK, etc.) for analysing each marker, and of colour or grey scale ranges for positive and negative objects.
   - Use of morphological filters in case they are needed to increase the efficiency of detection algorithms.
   - Detection of nuclear or cell edges.
   - Division of cell clusters in the images, if necessary.
   - Mathematical correction or extrapolation, if necessary.
   - Development of some algorithms by collaborating centres (University of Castilla La Mancha, Spain, and the Nalecz Institute of Biocybernetics and Biomedical Engineering, Warsaw, Poland).

7. Automated image analysis: The automated procedures developed in the present project will be based on automated procedures developed in our previous studies. The main features to take into account for the development of new procedures will be stain localisation (nuclear, membrane or cytoplasmic), and the presence of background and its stain intensity in image. In these studies the development of the image analysis procedures will be explained step-by-step. For the evaluation of the new markers included in the present project, the previous automated methods will be adapted using the image analysis techniques previously described in the part 6. Despite the large number of images produced (11 000), it will not take very long to analyse them. The image analysis procedures developed for each marker will be run consecutively, the analysis of 50 images taking approximately 20 min.
8. Compilation of patient data: Clinical and pathological data from patients enrolled in the study will be collected from electronic and paper medical records during the first year of the project. Clinical information related to the amount and localisation of DCs and other immune cells infiltrating the tumour and axillary lymph nodes will be collected during the second part of the third year of the project after quantifying them with image analysis procedures. Survival and relapse data will be collected during the last trimester of the study in order to have the longest follow-up. The date from last follow-up will be the date of the most recent visit to the hospital or primary care centre.

9. Analysis of results: The results will be of two types. A. Validation of image analysis procedures: Differences, reproducibility and concordance between manual and automated methods will be evaluated by the intraclass correlation coefficient, Student’s t test, Bland-Altman and Kaplan-Meier graphs and ROC curves. B. Immune response patterns: The evaluation of immune response of each case will be based on the mean of the evaluation of a total of four images: two images taken from each of the two cylinders of each case. The cases where one cylinder is lost due to the cut of samples or a bad immunostain will be discarded from the analysis. We have estimated a 10% of samples loss that have been included in the sample size calculation mentioned above.

Illness progression (relapse or not) and survival function (alive or dead) based on the immune infiltrates and the other clinicopathological variables will be evaluated by the Kaplan-Meier method and Cox regression to evaluate the association between the various markers and progression/survival of illness. The methodology for evaluating the relation of immune infiltrates with cancer progression on patient’s survival has been previously tested in our group’s previous studies. The statistics methods for this evaluation are described below.

The Kaplan-Meier method for estimating DFS and OS will be used. Two groups will always be compared using several cut-off points in the number of cells or in the level of molecular marker expression: mean, first quartile, second quartile, third quartile or the presence or absence of stained molecular markers.

The concentrations of cells and molecular markers will be compared between patients with and without metastasis. They will be analysed by the \( \chi^2 \) test when grouped as categories and by the Student t test when evaluated quantitatively. The same type of analysis will be performed for clinical variables depending on whether they are quantitative or categorical.

A Cox regression model will be fitted to compute the HR and its 95% CIs relapse/survival with each biomarker in the study. Those biomarkers with significance under 0.10 will be included in a multivariable Cox Regression model to obtain a relapse prediction model. Those clinical variables that could influence patients’ relapse will also be included in the multivariable Cox Regression model. Statistical analysis will be performed with the software IBM SPSS Statistics V.21.0.

Study limitations

Normally, a percentage of images cannot be analysed by automated methods because of poor staining. We have noticed that this happens more frequently in very old (pre-2000) samples or with immunohistochemical stains that are not widely used. However, in this project, the samples will have been obtained after 2000 and most of the markers involved are widely used in pathology departments. Under these conditions we would expect that a relatively low percentage of images would prove not possible to analyse.

ETHICS AND DISSEMINATION

Patients whose biopsies and clinical data will be used will provide signed informed consent. Results will be published in peer-reviewed journals.

The study guarantees the confidentiality of patients’ data. The data collected and generated will be included in two databases as a data-decoupling procedure to ensure adherence to data protection legislation and to maintain individual privacy. For this reason, different types of files will be created and maintained, each with a different password. Passwords will be changed periodically and will only be known to the person in charge.

The first file will be a demographic database in which each patient is assigned a unique code. The second file will include the patient codes linked to the information relevant to the study, including the results of the study and clinical variables for statistical analysis. The databases will remain stored on the renewably password-protected hard disk of a PC located in a locked office with controlled access. Results will be published in peer-reviewed journals and disseminated in international conferences and meetings related to digital pathology, immunology or cancer. It is expected that at least three articles will be published, two of which will be cover image analysis procedures and the third will present the clinical findings related to immune infiltrates and patient survival and relapse.

DISCUSSION

To understand cancer initiation and progression, it is necessary to focus on the cancer cell and on the tumour microenvironment from which the cancer cell obtains its nutrients, oxygen and signals for growth and development. The concept of the tumour microenvironment, developed from Paget’s ‘seed and soil’ theory, describes a dynamic network that includes the cancer cells, stromal tissue and the extracellular matrix that surrounds them.
Immune cells are an important class of stromal cells within the tumour microenvironment. Very few studies have evaluated the immune response in tumours and in lymph nodes simultaneously in the same patients. Moreover, with the development of more accurate methods for analysing immune infiltrates, it is becoming clearer that different infiltrating cell types have distinct prognostic and predictive significance. Several studies have demonstrated the presence of a heterogeneous population of immune cells in the breast tumour microenvironment, consisting mainly of T lymphocytes, cytoxic cells, tumour-associated macrophages, DCs, and regulatory cells. The types and amounts of these infiltrated immune cells are significantly associated with some clinical variables, such as tumour size, age, disease progression and survival, as well as the clinicopathological behaviour of the patient. Moreover, more than just the composition, distribution, architecture and functional articulation of the immune infiltrate, the immune context must be analysed to understand its significance and to derive more accurate prognostic and predictive biomarkers. This may involve analysing and evaluating the composition of the immune cells in the axillary lymph nodes.

A better understanding of the immune response, in particular that of DCs in patients with breast cancer, may help improve the design and application of therapeutic vaccines, making them more effective. In breast cancer, immunotherapy has been shown to prevent relapse, to improve survival and eliminate breast cancer in humans. Immunotherapy fits within the definition of personalised treatment, which is one of the two key technologies that is transforming pathology. The second of these technologies is digital pathology. Although the validation of these digital approaches has reaffirmed the accuracy of pathologists and the semiquantitative scoring methods they use, these new methods potentially require less pathologist time and resources, and produce a more objective assessment. There are two additional technologies that allow the in situ evaluation of a large number of molecular markers in a more rapid and homogeneous manner. The first is IHC, a cheap and accessible diagnostic technique used in daily clinical practice in pathology departments. Antibodies used in IHC are among the most frequently used tools in modern biomedical research, their popularity being reflected in the large number of immunohistochemical studies carried out over the past 20 years. The second is the TMA, which has become an established and valuable tool in cancer research, particularly in translational research and clinical trials. It allows resource-efficient use and high-throughput profiling of large numbers of tumours. The use of TMAs to assess biomarkers has become significantly more common over the past few years. Although the degree of correlation between TMAs and whole-tissue sections may not be considered ideal at the diagnostic level for individual patients, it is widely regarded as adequate in the research setting, where large samples are available for clinical trials or investigative studies.

In the present project, we expect that the use of TMAs and digital image analysis will enable us to identify differences in the patterns of immune response cells and specifically of DC infiltrates in the tumour and axillary lymph nodes between patients with and without metastasis in the axillary lymph nodes. We envisage being able to use this information not only as a source of prognostic factors but also to help improve the design and application of therapeutic vaccines, thereby making them more effective.

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Contributors CL as principal investigator, and ML as senior researcher, were involved in the conception of the project, the study design and drafting the manuscript. RB, JJ and TA evaluated and revised the patient biopsies and selected the cylinders for TMA construction. CL, MO and MoL compiled the pathological and clinical data of all patients enrolled in the study and created the database. AK, JB, GB and MG-R were involved in standardising image capture conditions and developing the image analysis procedures. MdMF-C, CC and GO captured images and applied image analysis procedures to them. CL, ML and MTS developed the final database and statistically analysed the results. CL drafted the first version of the manuscript. All authors read and approved the final version of the manuscript.

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Competing interests CL, RB, JJ, MTS, TA, MoL, JB and ML are members of the Oncological Pathology and Bioinformatics Research Group of the Hospital de Tortosa Verge de la Cinta (Pere Virgili Health Research Institute, Spain). CC is a PhD student associated with this study and GO works in the Molecular Biology and Research section of the Hospital de Tortosa. MO was a gynaecologist at the Hospital de Tortosa. AK is an engineer at the Nalecz Institute of Biocybernetics and Biomedical Engineering, Warsaw, Poland. MG-R is a specialist in telepathology at Hospital de Jerez, Spain. GB and
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