Case Report

Rosai–Dorfman Disease: Breast Involvement—Case Report and Literature Review

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Abstract: Background and objectives: Rosai–Dorfman disease (RDD) is a type of histiocytosis that usually appears in young adults or children as bilateral cervical lymphadenopathy, but extranodal involvement is not uncommon. Although the pathogenesis is not entirely elucidated, recent studies showed a possible neoplastic process. Materials and methods: Our manuscript presents a rare case of Rosai–Dorfman disease of the breast, the management of this rare case, and a literature review. There are few cases reported of RDD of the breast (around 90 globally reported cases); the data is poor, and the management not yet standardized for these cases. The case reported here shows the importance of correct breast investigation, breast imaging, and ultrasound-guided biopsy that provided an accurate diagnosis and guided further management. Results: Although RDD of the breast was rarely presented as bilateral disease in other case reports, our case showed bilateral breast disease with the suspicion of breast cancer on imaging. Pathology and immunohistochemistry were of critical importance and showed a specific pattern for histiocytosis. A multidisciplinary approach was taken into consideration for these cases in order to establish the approach. Some patients underwent surgery, but watchful waiting and close follow-up were the preferred approach. Conclusions: RDD of the breast is a rare form of histiocytosis, with fewer than 100 globally published cases. Although the management of this disease is not established yet by guidelines, a follow-up approach should be enough for these patients, and surgery might be overtreatment. Mortality from RDD is very low due to comorbidities. A multidisciplinary team decision is important, and abstinence might significantly benefit these patients.

Keywords: Rosai–Dorfman disease; Rosai–Dorfman of breast; breast pathology; histiocytosis

1. Introduction

Rosai–Dorfman disease (RDD) is a histiocytosis that involves usually cervical lymph nodes in children and young adults. However, extranodal involvement is not uncommon, often raising diagnostic and therapeutic difficulties. It was described by a French pathologist, Pierre Destombes, as early as 1965, and was then thought to be a lipid storage disease occurring as a consequence of inflammation [1]; four years later, Rosai and Dorfman correctly identified the key roles of histiocytes in the pathogenesis of disease, most often presenting as bilateral cervical painless lymphadenopathies, and communicated 34 similar cases with voluminous lymphadenopathy [2–4].

Currently, it belongs to R-group histiocytosis according to the revised classification of The Histiocyte Society, which includes familial RDD, classical (nodal) RDD, extranodal...
RDD, neoplasia-associated RDD, immune disease-associated RDD, and other miscellaneous types of histiocytosis [5].

Extranodal RDD is encountered in over 40% of patients and usually occurs concomitant with nodal disease. The most frequently involved extranodal localizations cited in the literature are the nasal cavity, skin, orbits, bone and central nervous system, but also skeletal muscle, subcutaneous tissue, heart, or thyroid. Breast localization is rare and often causes difficulties in differential diagnosis with breast malignancies. Other localizations are also cited, such as in the eyes, gastrointestinal tract, or uterine cervix [6,7].

The etiology of the disease is still unknown, and its natural history is marked by long phases of remission and relapse. Some theories suggest immune regulation disorder or infection etiology such as the varicella zoster virus, herpes virus, Epstein–Barr virus, cytomegalovirus or HIV [8,9].

Clinically, the classical nodal disease is painless, and examination reveals bilateral cervical lymphadenopathy in most cases. Associated symptoms can be fever and night sweats or weight loss. Apart from cervical lymphadenopathy, other groups of lymph nodes can also be involved (axillary, inguinal, mediastinal, or rarely retroperitoneal) [10,11].

Apart from the classical nodal RDD, extranodal RDD can involve a large variety of localization, with symptoms according to the involved organ (double vision or orbital pain, nasal obstruction or epistaxis, oral pain, dyspnea, cough, abdominal pain, hematochezia, hematuria, bone pain, skin changes, headaches, seizures, or other neurologic manifestations) [12].

RDD on laboratory evaluation is associated with a large number of neutrophil cells on full blood count with differential polyclonal hypergammaglobulinemia on serum immunoglobulins testing and high erythrocyte sedimentation rate. Other laboratory tests to be considered for differential diagnosis or associated disease identification are antinuclear antibodies, HLA-B27, or autoimmune lymphoproliferative syndrome markers, complete metabolic panel, blood smear or bone marrow aspirate, and lumbar puncture [12].

Pathology examination for nodal RDD reveals large histiocytes with pale cytoplasm and large hypochromatic nucleus and sinus expansion. Immunohistochemistry shows nuclear and cytoplasmic S-100 and CD68 positivity, and sometimes CD163 and CD14 positivity. RDD is differentiated by Langerhans cell histiocytosis by CD1a and CD207 negativity. These characteristics help differentiate RDD from Langerhans cell histiocytosis and Erdheim–Chester disease. Emperipolesis (intact leukocytes in the cytoplasm of histiocytes) is a useful marker but is not a mandatory finding for diagnosis, as it is sometimes missed on evaluation because of the focal presence, usually in extranodal sites, and it can also be found in other pathologies, namely, Erdheim–Chester disease, juvenile xanthogranuloma, or malignant histiocytosis [12].

Lymphoma and infectious diseases such as tuberculosis are the most common clinical entities that can mimic RDD, especially in developing countries. Moreover, the disease appears more often in male children or young adults. The unusual presentation of breast lumps in premenopausal women can easily be considered to be a breast malignancy, this being atypical for RDD [13].

The treatment strategies for RDD include an expectative approach and monitoring, corticosteroids (usually prednisone or dexamethasone), surgery for unifocal and/or symptomatic extranodal disease, radiotherapy (with a dose of 30–50 Gy), chemotherapy (Vinca alkaloids, methotrexate, cladribine), or immunomodulatory therapy (thalidomide, lenalidomide, rituximab, imatinib mesylate) [12,14,15].

Although the clinical course is unpredictable and marked by long phases of remission with periods of relapse and worsening symptoms, the disease seems to be self-limiting in many cases in which observation is reasonable, with 20%–50% of patients with nodal or cutaneous RDD experiencing spontaneous remission. However, evidence is scarce because of the lack of randomized trials. The overall outcome and prognosis of RDD are usually good, and treatment is necessary in selected cases with important clinical
signs such as vital-organ compression and airway obstruction due to the impressive node enlargement [15].

Outcome is favorable in the majority of cases, especially for nodal or cutaneous diseases; death as a direct result of RDD is rare, cited between 7% in one of the largest series of patients with RDD published and 12% in a comprehensive review published in 2002 [15,16].

We analyzed extranodal RDD involving the breast in our review because it is a rare condition, currently described in the literature in only about 90 cases, with non-standardized management because of the rarity of the disease and because existing evidence regarding the management of this localization is still scarce.

2. Objective

The principal objective of this comprehensive review and case report is to bring new insights and to systematize the existing data in the literature about this rare localization of RDD, helping to increase the accuracy of diagnosis and adequacy of treatment of this rare condition of the breast.

3. Case Presentation

We present the case of a 63-year-old woman with a personal history of renal neoplasia (transitional cell carcinoma surgically treated eight years ago, free of disease at present) and systemic histiocytosis (bone located), and a family history of lung carcinoma (father) and thyroid carcinoma (daughter). Systemic histiocytosis was diagnosed six years ago on a surgical specimen from a tibial bone tumor. No pathology results were available. Thereafter, she did not have any other manifestations of disease until breast lumps were diagnosed on screening mammogram. She was not under any treatment for her systemic histiocytosis.

She presented with bilateral breast lumps for investigation and management. Clinical examination revealed bilateral breast lumps on the right side in the axillary tail of the right breast, a palpable lump of about 2.5 cm with irregular borders, not infiltrating the skin or muscles, and on the left side, at the border of inferior quadrants, a 1.5 cm lump with similar characteristics; no palpable suspicious axillary lymph nodes were clinically identified. No other suspicious lesions located outside the breasts were identified.

Mammography revealed bilateral suspicious findings: an opacity on the right side towards axillary tail with diameters 2.0/1.6 cm, slightly irregular, and on the left side a similar image of 1.3/1.4 cm, distributed at the border between inferior quadrants (Figures 1 and 2). Ultrasound imaging confirmed the two above-described lesions. They were both scored BIRADS 4a. There were no suspicious axillary lymph nodes visualized on imaging.

Bilateral ultrasound-guided 14-gauge core biopsies of the breast lumps were undertaken. Pathology exam findings (haematoxylin–eosin and immunohistochemistry) revealed histiocyte aggregates, emperipolesis, histiocytes diffuse positive for S100 protein, CD1a, focally expressing CD68, negative for CD20 and CD3. Pathology findings were consistent with the diagnosis of bilateral Rosai–Dorfman disease of the breast.
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Figure 3. Rosai–Dorfman disease of the breast—haematoxylin–eosin (20×).

Figure 4. Rosai–Dorfman disease of the breast: immunohistochemistry—S100 diffuse positive histiocytes (20×).
The multidisciplinary team meeting decision was to continue surveillance with imaging in the absence of any medical or surgical treatment.

The patient repeated ultrasound assessment at 6 months and mammogram at one year follow-up. The breast lumps disappeared at 6-month follow-up without any treatment.

The particularity of our case is the early diagnosis after suspicious imaging using an ultrasound-guided core biopsy in a patient with a suggestive personal history of RDD and conservative management of the breast tumour with follow-up imaging, avoiding unnecessary surgery.

4. Materials and Methods

For the present comprehensive review, we selected all published original articles communicating confirmed cases on pathology with extranodal Rosai–Dorfman disease located in the breast and/or axilla.

A comprehensive review of the literature was undertaken; eligibility criteria circumscribed all original articles reporting on extranodal Rosai–Dorfman disease of the breast published in the literature, including case reports and case series. All selected papers' reported cases confirmed on pathology with breast and/or axillary localization.

A literature search was undertaken using as keywords “Rosai–Dorfman disease”, “histiocytosis”, and “breast”. The PubMed, Embase, and Scopus databases were searched. Articles published in English until April 2021 were selected, although other languages were not excluded (Spanish, Dutch, Chinese). We found 55 articles; of these, 7 articles were excluded after reading the abstract, and 6 articles after reading the full text. The excluded articles were not relevant to our study because they were review papers or lacking original case reports, or RDD was not located in the breast. We selected 42 articles after reading the abstract and full text (selection algorithm in Figure 7). Selected articles are systematized in Table 1 [17–58]. The strength of evidence-based data was low because of the rarity of RDD localization and the small number of patients; only case reports, case series, and retrospective trials were found on extensive search, and no prospective or randomized data were identified.
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| Total abstracts selected after first search: 55 articles |
| Articles excluded by abstract: 7 articles |
| Full read articles: 48 articles |
| Studies that did not match criteria: 6 articles |
| Final selected articles: 42 articles |

Figure 7. Algorithm of article selection.
| Author                  | No of pts | Age  | Breast Side | Breast Localization | Axillary | Diagnosis | Pathology | Management | FU | Gender | Observations                  |
|------------------------|-----------|------|-------------|---------------------|----------|-----------|-----------|------------|-----|--------|-------------------------------|
| Battle 2021 [17]       | 1         | 49   | RB          | Multifocal, UIQ     | no       | 1,2,3     | 1         | 1          | N/A | F      | N/A                           |
| Delaney 2017 [18]      | 1         | 63   | RB          | UOQ                 | no       | 1,2,3     | 1         | 1          | 1   | F      | N/A                           |
| Liu 2018 [19]          | 4         | 46–68| N/A         | N/A                 | no       | 1,2,3     | 1         | 1          | N/A | F      | N/A                           |
| Hoffmann 2019 [20]     | 22        | median 54 (range 37–71) | RB, LB | N/A | no | 1,2,3 | 12 | 4 cases—1; 18 cases—2; (1 case mastect) | 4 cases—1; 2 cases—2 | 18—F, 4—M | N/A |
| Chen 2016 [21]         | 12        | median 37 (range 28–47) | 7—RB, 5—LB | N/A | yes—1 no—11 | 1,2,3 | 2 | N/A | N/A | 12—F | Article in chinese |
| El-Attrache 2018 [22]  | 1         | 55   | RB          | LQ                  | no       | 1,3       | 1,2       | 2          | 2   | F      | Breast recurrence of RDD     |
| Shetty 2020 [23]       | 4         | median 58 (range 43–69) | N/A | N/A | yes—2 no—2 | 3—1,2 | 1,2 | 2 | 3 cases—1; 2 cases—3 1 case—N/A | F | N/A |
| Shin 2020 [24]         | 1         | 54   | RB          | OUQ                 | no       | 1,2,3     | 1,2       | 1          | 1   | F      | N/A                           |
| de Mello Tucundova 2017 [25] | 1 | 55 | LB | N/A | no | 1,2,3 | 1,2 | 2 | N/A | F | N/A |
| Jorns 2017 [26]        | 1         | N/A  | LB          | N/A                 | no       | 1,2,3     | 1,2       | 1          | 1   | F      | No recurrence                 |
| Goldbach 2019 [27]     | 1         | 44   | RB          | UIQ                 | no       | 1,2,3     | 1,2       | 2          | 1   | F      | Recurrence at 6 months       |
| Parkin 2015 [28]       | 1         | 56   | LB          | N/A                 | no       | 1,2,3     | 1,2       | 1          | 1   | F      | N/A                           |
| Ciurea 2016 [29]       | 1         | 59   | RB          | UIQ                 | no       | 1,2,3     | 2         | 1          | 1   | F      | Hyperpigmentation of tegument |
| Tenny 2011 [30]        | 1         | 64   | RB          | N/A                 | yes      | 1,2,3     | 1,2       | 2          | 2   | F      | Multiple distant recurrence at 6 months |
| Morkowski 2010 [31]    | 3         | median 51 (range 39–62) | 2 LB, 1 RB, 2 USQ | N/A | no | 1,2,3 | 1,2 | 2 | 1 | F | No recurrence after excision |
| Author         | No of pts | Age   | Breast Side | Breast Localization | Axillary | Diagnosis | Pathology Mangement | FUp | Gender | Observations                       |
|---------------|-----------|-------|-------------|---------------------|----------|-----------|---------------------|-----|--------|-----------------------------------|
| Vaidya 2020   | 1         | 43    | LB          | UOQ                 | no       | 1,2,3     | 1,2                 | 2  | 1      | F                                 |
| Simmons 2016  | 1         | 41    | RB          | Multifocal          | no       | 1,2,3     | 1,2                 | 2  | 1      | F                                 |
| Zhou 2016     | 1         | 71    | LB          | Multifocal          | no       | 1,2,3     | 1,2                 | 2  | 1      | F                                 |
| Green 1997    | 1         | median 46 (range 15–84) | 4 RB, 2 LB axillary | 2—bilateral | yes—1 no—6 | 1,2,3 | 1,2 | 6—2 1—1 | N/A | F | N/A |
| Cha 2012      | 1         | 62    | RB          | LOQ                 | no       | 1,2,3     | 1,2                 | 2  | 1      | F                                 |
| da Silva 2007 | 1         | 50    | LB          | UOQ                 | no       | 1,2,3     | 1,2                 | 2  | 1      | F                                 |
| Ng 2000       | 2         | median 50 (47–58) | 4 RB | UOQ | no | 1,FNA 1,2 | 1,2 | 2 | 1 | F | No recurrence at 1 and 6 months FU |
| Moyon 2020    | 1         | 29    | N/A         | N/A                 | no       | 1,2,3     | 1,2                 | 1  | N/A    | F                                 |
| Bansal 2010   | 1         | 35    | RB          | LOQ                 | no       | 1, FNA 1,2 | 1,2 | 1 | 1 | M | No recurrence at 18 Mo FU |
| Fu 2012       | 1         | 78    | RB          | LOQ                 | no       | PET/CT scan | 1,2 | 2 | N/A | F | N/A |
| Mantilla 2016 | 2         | 63    | LB          | 1 multicentric      | no       | 1,2,3     | 1,2                 | 1  | 1      | F | At 3 years FU—subcutaneous soft mass |
| Krbanevic 2021| 1         | 50    | Bilateral   | N/A                 | no       | 1,3       | 1,1                 | 1  | N/A    | F                                 |
| Wu YC 2010    | 1         | 33    | RB          | UOQ                 | no       | 1,3       | 2,3                 | 1  | F      | No recurrence at 2 years FU      |
| Mac-Moune Lai | 1         | 34    | LB          | LIQ                 | no       | 1,2       | 2                   | 2  | 1      | M | No recurrence at 3 months FU     |
| Wang 1997     | 1         | 35    | LB          | N/A                 | no       | 1,2,3     | 2                   | 2  | 2      | F | Recurrent breast tumor            |
| Gwin 2011     | 1         | 68    | Bilateral   | N/A                 | no       | 1,2,3     | 1,2                 | 1  | N/A    | F                                 |
| Noordzij 2011 | 1         | 75    | RB          | Multifocal          | no       | 1,3       | 1,2                 | 1  | N/A    | F | Article in Dutch                  |
| Author                | No of pts | Age | Breast Side | Breast Localization | Axillary | Diagnosis | Pathology | Mangement | FUp | Gender | Observations                  |
|-----------------------|-----------|-----|-------------|---------------------|----------|-----------|-----------|-----------|-----|--------|-------------------------------|
| Pham 2005 [49]        | 1         | 53  | LB          | LIQ no 1,2,3        | 1,2,3    | 1,2       | 1         | N/A       | F   | N/A    |                               |
| Elshikh 2020 [50]     | 3         | 60  | LB          | N/A no 1,2          | 1,2      | 1         | 1         | N/A       | F   | N/A    |                               |
| Hammond 1996 [51]     | 1         | 67  | RB          | UOQ no 2,3         | 1,2,3    | 1,2       | 2         | 1         | F   | N/A    | No recurrence at 6 Mo FU     |
| Kuzmiak 2003 [52]     | 1         | 30  | RB          | UOQ, Multifocal no 1,2,3 | 1,2,3 | 1,2       | 2         | N/A       | F   | N/A    |                               |
| Baladandapani 2012 [53]| 1       | 59  | LB          | Multifocal, UQ no 1,2,3 | 1,2,3 | 1,2       | 2         | N/A       | M   | N/A    |                               |
| Hummel 1999 [54]      | 1         | 52  | LB          | UIQ no 1           | 1        | 1,2       | 2         | 1         | F   | N/A    | No recurrence at 11 Mo FU    |
| Dahlgren 2008 [55]    | 1         | 64  | Bilateral   | Bilateral no 1,2,3  | 1,2,3    | 1,2       | 2         | N/A       | F   | N/A    |                               |
| Dias Perera 2007 [56] | 1         | 23  | LB          | N/A no 1,2         | 1,2      | 2         | 2         | N/A       | M   | N/A    |                               |
| Picon-Coronel 2010 [57]| 1       | 67  | N/A         | N/A no 1,2         | 1,2      | 1,2       | 2         | N/A       | F   | Aticle in spanish            |
| Perez-Guillermo 1993 [58]| 1      | 71  | Bilateral   | UIQ left, LIQ right no 1,2,3 | 1,2,3 | 1,2       | 2         | N/A       | F   | N/A    |                               |

Abbreviations: F- Female, M- Male, LB—left breast, RB—right breast, N/A not available, FU—follow up, UOQ—upper outer quadrant, UQ—upper quadrants, UIQ—upper inner quadrant, LOQ—lower outer quadrant, LIQ—lower inner quadrant, FNA—fine needle aspiration. Diagnosis—1—US; 2—Mammogram; 3—Biopsy. Pathology—1—emperipolesis, 2—S100 positive, 3—CD46 positive. Management—1—expectant, 2—surgery, 3—steroids. Follow-up—FU—1—no progress or recurrence, 2—recurrence, 3—death.
5. Results

A total of 92 patients with a diagnosis of Rosai–Dorfman disease of the breast on pathology were selected and analysed. Most literature communications were case reports or case series. The median aged was 55 years old (15–84), while gender distribution was dominated by females (10 males to 82 females).

Imaging was performed in most patients, usually using ultrasound and/or mammogram, often followed by imaging-guided core biopsy. Axillary involvement was described in 5/92 patients (5.4%), which confused the clinical picture even more, orientating the clinical suspicion towards malignancy.

Multicentric appearance at diagnosis was rather rare, communicated in 7 out of 78 patients (9.0%). Bilateral breast involvement with data provided in the selected papers was described in 5/78 patients for (6.4%). Recurrent disease was uncommon, cited in 6/64 patients (9.4%) (for 28 patients, there was no follow-up reported). There was no significant association between axillary or bilateral breast involvement and systemic disease manifestations.

Pathology diagnosis is critically important in breast abnormalities. New pathology categories based on immunohistochemistry are currently researched [59]. RDD of the breast was definitively diagnosed on pathology either on biopsy or on surgical specimen. Of the patients, 55/92 (59.8%) underwent core-needle biopsy, while the rest had a diagnosis on the surgical specimen. Pathology diagnosis was conducted by the identification of large histiocytes with a round nucleus containing vesicular chromatin with a couple of nucleoli and pale cytoplasm; the typical finding was emperipolysis (from Greek: em—inside, peri—around, polemai—wander about), a phenomenon consisting of the presence of lymphocytes in the cytoplasm of other cells, usually histiocytes; it is defined as the penetration of one living cell by another cell that remains intact, distinct from phagocytosis. [60]

Pathology diagnosis was eased by immunohistochemistry staining of S-100 protein and CD-68 in most cases, characteristic for antigen-presenting or Langerhans cells. Most presented cases of RDD of the breast did not have molecular study evaluation or assessed BRAF mutations.

More than half of the patients (56 out of 92 patients—60.9%) underwent surgery, usually lumpectomy or even mastectomy [20]. The rest were approached expectantly; steroids were administered in a minority of patients.

Follow-up information varied from lost to follow-up to 3–12 months, while a few studies cited 2 or 3 years follow-up [42,44]. The course of the breast RD disease was usually uneventful with spontaneous remission in the majority of patients and with no recurrence in patients that underwent surgery. Breast recurrence was usually the exception during the course of disease [24,30], while fatalities during follow-ups were also very rare and due to other causes (acute leukemia, respiratory failure [23]).

6. Discussion

Our study sheds more light in the rare pathology of RDD of the breast and systematises existing data in the literature regarding the diagnosis and management of this condition. Because of the paucity of literature data, the differential diagnosis and management of RDD of the breast are still debated, with the current data reflecting the difficulty of differential diagnosis and frequent surgical overtreatment.

Breast involvement in Rosai–Dorfman disease is a rare occurrence, but it can be mistakenly considered to be a malignant condition on clinical examination, imaging, or even pathology [30]. The involvement of the axilla can induce even more confusion for the clinician. Most cases communicated in the literature were postmenopausal women diagnosed after 50 years of age using common breast imaging tools such as ultrasound and mammogram. A personal history of Rosai–Dorfman disease helped in raising suspicions of extranodal RDD. Final diagnosis was performed on pathology on either core needle biopsy or excisional specimen. Emperipolysis on HE staining and staining for CD-68 and S-100 on immunohistochemistry were cardinal features for diagnosis.
Differential diagnosis on pathology is usually performed with inflammatory myofibroblastic tumours, granulomatous mastitis, IgG4-sclerosing mastitis, or breast lymphoma with plasmocytic differentiation.

In our patient, the diagnosis of RDD of breast was eased by the personal history of histiocytosis, although imaging was suspicious of malignancy. Core needle biopsy limited the morbidity of an excisional biopsy and pathology diagnosis of RDD of the breast spared the patient any surgical treatment, leading to spontaneous disappearance of bilateral breast nodules at one year follow-up.

Treatment of this localization of RDD was not necessary in most of the cases. However, due to paucity of literature data, many patients underwent surgical excision or lumpectomy, with diagnostic and curative intent. Only one case of radical surgery (mastectomy) was found in the literature [20].

The outcome of RDD breast localization seems to be very good even without any intervention. The occasional progression or recurrence of the disease was cited in the literature, and mortality was exceptionally communicated, usually due to other comorbidities.

7. Conclusions

RDD of the breast is a rare localization that can confound diagnosis and lead to surgical overtreatment. Clinicians should be aware of its existence and raise suspicion, especially in patients with personal history of RDD. Core needle biopsy is needed for definitive diagnosis; clinicians should effectively communicate with pathologists for a timely and correct diagnosis. Pathology diagnosis of emperipolesis and positive staining on immunohistochemistry for S100 protein and CD-68 is of utmost importance. Until more evidence-based data are available, a conservative approach with abstention from any kind of surgery and clinical follow-up would spare the patients the risks of unnecessary treatment.

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