Idiopathic Granulomatous Lobular Mastitis – Report of 43 Cases from Iran; Introducing a Preliminary Clinical Practice Guideline

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Keywords
Idiopathic granulomatous lobular mastitis · Multidisciplinary approach · Treatment strategies · Clinical practice guidelines · Iran · Milk extravasation · Breast-feeding · Relapse

Summary
Background: We aimed to report a large series of idiopathic granulomatous lobular mastitis (IGLM) from Iran and sketch preliminary clinical practice guidelines (CPG) for approaching an inflammatory breast mass. Patients and Methods: In a retrospective records review, 43 consecutive IGLM cases were studied. Data on baseline, clinical, imaging, and pathologic characteristics were collected. Results: The mean age of the women was 33.5 years. All but 1 were married and had given birth. 16% had a cancer-like presentation. Inflammatory signs, architectural distortion, and a nodular pattern were the most common findings clinically, mammographically and ultrasonographically, respectively. 29.5% of the pathological reports indicated necrosis which was more common in younger subjects (p = 0.016); microabscesses were associated with a shorter lactation course (p = 0.006). Corticosteroids had been used as the initial treatment modality in 51%, immunosuppressive agents had not been administered, and a 16% relapse rate was recorded. We recognized the need for a multidisciplinary approach covering radiology, oncology, and surgery to best handle diagnostic and therapeutic issues and manage relevant infections as well as the major differential diagnosis, i.e. malignancy. Conclusion: We hypothesized that a shorter lactation period may cause more milk stasis and extravasation and be contributory to IGLM. CPGs are needed to incorporate the needed multidisciplinary approach and to standardize IGLM care. We present one such guideline.

Schlüsselwörter
Idiopathische granulomatöse lobuläre Mastitis · Multidisziplinäres Vorgehen · Behandlungsstrategien · Klinische Praxisrichtlinien · Iran · Milchaustritt ins Gewebe · Stillen · Rezidiv

Zusammenfassung
**Introduction**

Idiopathic granulomatous lobular mastitis (IGLM) is a rare relapsing inflammatory pseudotumor which is commonly mistaken for malignancy [1, 2]. It presents as an ill-defined indurated and relatively large mass, commonly in the upper outer quadrant or subareolar zone of the breast, typically in the reproductive post-childbearing period [3]. The mass is tender and may be associated with nipple retraction, peau d’orange-like changes, and an overlying erythematous and/or ulcerated skin. These features make the differential diagnosis challenging [3, 4]. Ultrasonographic, mammographic, and even magnetic resonance imaging findings are not pathognomonic. Fine needle aspiration cytology is practiced, but in nearly all cases core needle (or open) biopsy becomes mandatory [4, 5].

There is no consensus about the management of IGLM. The relapsing nature is sometimes frustrating and the cosmetic sequelae of surgical excision are unwelcome. The aim of the current study is to report a relatively large series of IGLM from Iran and to sketch a preliminary clinical guideline for approaching an inflammatory breast mass.

**Patients and Methods**

In a retrospective study, records of 43 cases of confirmed IGLM were reviewed (Breast Cancer Clinic, Cancer Institute, Tehran; 2006–2012). Data on baseline, clinical, imaging, and pathologic characteristics were collected. The setting was a cancer referral center in the Capital Tehran, and the patients had a minimum follow-up of 6 months (median 16 months). Meticulous breast examination including the axillary region had been performed. The mentioned signs in the clinical records were collected. Mammography had been routinely carried out for the cases 35 years of age or older. An ultrasonography and fine needle aspiration biopsy had been variably performed, but the definitive diagnosis had been reached either through 14-gauge core needle biopsy or surgical excision and pathologic confirmation. In cases of suspicion for tuberculosis and bacterial infection, Ziehl-Neelsen and gram staining had been carried out. A tuberculin skin test had been ordered likewise. We encountered a single case of tuberculosis which was excluded from the series of 44 cases. Non-parametric statistical testing (Mann-Whitney U test) was performed to explore the relationship between relapse and pathologic features with potential baseline and clinical course factors. Records were coded and kept anonymous.

**Results**

The mean age of the patients was 33.5 years (range 24–49 years), excluding a single woman of 55 years of age. 92% of the patients were in the reproductive period, and 8% were in the perimenopause (> 45 years of age). The mean of the reported age at menarche was 14 years (range 12–16 years). All but 1 case were married, had given birth (median 1 child, range 0–5 children), and had a history of breastfeeding (mean 30 months, range 0–120 months). None of the patients had a positive family history of breast cancer. The right breast was affected in 44% and the left breast in 56% (2 cases were bilateral). The median period of symptom-to-referral was 2 months, but the range was up to 4 years.

The cases presented with a variety of sign and symptom complexes (table 1). The presentation in 7 (16%) cases was suggestive of cancer, and 78% had a history of empiric antibiotic treatment (oral cloxacilin or cephalaxin for 10 days). In those who had had a mammogram (age > 35 years), architectural distortion, nodular lesions, and asymmetric density were reported in descending order. Sonography was abnormal in 95% of cases; a nodular pattern and (large or small) hypoechoic spaces were reported (table 1). A biopsy was taken within the first month of referral in almost all cases (open biopsy 43.5%, core biopsy 56.5%). The diagnosis was made by detection of granulomatous lesions (aggregates of epithelioid cells) in a ductolobular configuration. It was notable that more than half of the pathological reports documented (micro)abscesses, and in a third of all patients necrosis was observed. 45% of cases had a white blood cell count of above 10,000/mcl.

A total of 22 (51%) cases were initially treated with systemic steroids, and the remaining patients underwent surgical resection as the initial modality. 90% of the cases were symptom-free at the last follow-up, and in the remaining 4 cases the inflammation had subsided significantly. 5 (11.5%) cases of the steroid-treated group needed later surgical intervention, in some cases despite repeated courses of steroids. Of the 16% patients who had recurrences, one third was complicated by fistula formation necessitating surgical intervention.

**Table 1. Clinical, imaging, and pathologic findings of idiopathic granulomatous lobular mastitis in our series (figures rounded; cases may have been listed in more than 1 category)**

<table>
<thead>
<tr>
<th>Clinical presentation</th>
<th>Mammographic findings (39 cases)</th>
<th>Sonographic findings (39 cases)</th>
<th>Pathologic findings</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mass 57%</td>
<td>nodular lesion 21.5%</td>
<td>nodular pattern 79.5%</td>
<td>multinucleated giant cells 67% (micro)abscess formation 53%</td>
</tr>
<tr>
<td>Suggestive for malignancy 16%*</td>
<td>architectural distortion 28.5%</td>
<td>hypoechoic spaces 28%</td>
<td>mononuclear infiltration 79.5%</td>
</tr>
<tr>
<td>Inflammatory signs 34%</td>
<td>asymmetric density 14%</td>
<td></td>
<td>neutrophilic infiltration 57.5%</td>
</tr>
<tr>
<td>Fistula 9%</td>
<td></td>
<td></td>
<td>necrosis 29.5%</td>
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<tr>
<td></td>
<td></td>
<td></td>
<td>leukocytosis 45%</td>
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</tbody>
</table>

*aIncluding induration, skin retraction, ill-defined margins.
Notably, necrosis was more frequently reported in younger patients (mean 29 vs. 34 years; p = 0.016). Patients whose pathological reports indicated microabscess formation had a shorter history of lactation (19 vs. 37 months; p = 0.006). Relapse was not correlated with the method of biopsy, history of antibiotic treatment, mammographic and ultrasonographic findings, age, and gynecologic history (all p > 0.05).

**Discussion**

Recently, several series of IGLM were published in the international literature [6–9]. It is not known whether there is a real epidemic of the condition, or simple awareness and better reporting have caused this outbreak. We know that in recent decades the incidence of autoimmune diseases has been on the rise [10]. In the case of IGLM, changes in reproductive and lactation behavior might have contributed to this (fig. 1).

The characteristics of our cases were generally similar to those of previous reports. For instance, almost all patients were of a reproductive and childbearing age, and the mean age was consistent with previous reports [9, 11]; only 1 of our cases was older than 50 years. All but 2 of our patients had a history of pregnancies and breastfeeding. This is compatible with the fact that IGLM mostly happens in the first few years following pregnancy or in those with a history of contraceptive use [12].

Clinical, mammographic, and sonographic findings (table 1) emphasize that through clinical and imaging studies one cannot differentiate IGLM from breast cancer [13, 14]. Mammographic findings in our series did not show the most characteristics signs of cancer, i.e. irregular or branching calcification [15], but architectural distortion and nodular lesions were seen. We documented relatively fewer asymmetric densities [16]. Sonography findings were similar to previous reports [16] but were also not characteristic as they included hypoechoic spaces and nodular lesions both of which are frequently observed in malignancy as well; however, cancerous lesions are less well-defined and more heterogeneous.

The implication is that we have to include IGLM as an important entity in the differential diagnosis of breast cancer, at least in referral centers, in order to prevent unnecessary wide excision. The broader list covers cystic neutrophilic granulomatous mastitis (in which the gram-positive bacillus Corynebacterium is involved), tuberculous mastitis, sarcoidosis, and Wegener’s disease. Among the patients referred to us, a single tuberculous case was diagnosed and successfully treated.

It is noteworthy that 3 of our cases had coincident or previous erythema nodosum. The association of IGLM with other autoimmune phenomena has already been well described [11], and this emphasizes the necessity for both a systemic survey and a multidisciplinary approach in the management of this condition [13]. Due to the retrospective nature of this study, we did not have access to enough ordered lab tests for parameters such as erythrocyte sedimentation rate, C-reactive protein, rheumatoid factor, and antinuclear antibody [11]. However, the common presence of leukocytosis in our series points to the systemic nature of the condition, and we acknowledge that the patients with suspected IGLM require a serologic survey (table 2 and fig. 2). This is relevant for both the identification of coincidental rheumatologic disease and the monitoring of disease activity and response to treatment.

In histopathology, non-necrotizing granulomas (along with multinucleated giant cells) and ‘lobulocentric’ [17] arrange-

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**Table 2. Preliminary Clinical Practice Guidelines for Approaching Inflammatory Breast (Pseudo)Tumors**

<table>
<thead>
<tr>
<th>Notes</th>
<th>Management principles</th>
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<tbody>
<tr>
<td>i) IGLM is exceptional in that its diagnosis and management may require participation from a multitude of specialties; namely, infectious disease, radiology, pathology, rheumatology, oncology, surgery (general and reconstructive), and obstetrics! ii) Despite the acknowledgement of this feature [13, 28, 29], the multidisciplinary approach is inadequately practiced. iii) This preliminary guideline intends to formalize management principles and to incorporate multidisciplinary participation.</td>
<td>Figure 2 presents the typical approach.</td>
</tr>
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<td>iii) IGLM is an inflammatory ‘pseudotumor’, and surgical excision should not be the initial choice [1].</td>
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<td>iv) Do not discontinue systemic steroids prematurely due to the relapsing nature of the disease, and ensure full remission through imaging modalities such as sonography which is not diagnostic but more sensitive than clinical examination for follow-up [1].</td>
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<tr>
<td>v) Avoid longer than 6 months systemic steroids due to medical complications and as there are effective alternatives such as cytotoxic agents like methotrexate [30].</td>
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**Fig. 1. Hypothetical pathogenesis of idiopathic (‘milk-induced autoimmune’) granulomatous lobular mastitis.**

- Breast tissue hypertrophy & fragility (caused by hyperprolactinemia, oral contraceptives, pregnancy, lactation)
- Breast micro trauma e.g. nursing & milk stasis (abortive lactation/premature weaning) & interstitial extravasation
- Induction of autoimmunity (mediated by Corynebacterium) & relapses (autoimmune disease-associated)

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ment (with intact ductolobular architecture) are pathognomonic. However, this is not always the case; in our series necrosis was present in a sizable proportion of the patients (29.5%).

Granulomas are not characteristic per se, and in order to differentiate the aforementioned entities, ancillary tests are required. One may state that IGLM clinically resembles cancer, and pathologically tuberculous mastitis. In the case of tuberculous mastitis, there is more fibrosis and eosinophils, and (caseation) necrosis can be seen [6, 18]. In IGLM, however, granulomas are admixed with neutrophils and plasma cells. This is a feature of the IGLM histopathology, i.e. ‘mixed’ inflammatory infiltration (table 1). Some reports still link IGLM to Corynebacterium infection [19–21]; however, as was shown in our cases, in IGLM mononuclear cells are comparatively more common than neutrophils (table 1) and vice versa. It is noteworthy that in none of our cases calcification had been reported, and calcification has remained as a major differentiating factor between cancer and IGLM.

More frequent observations of necrosis in younger cases emphasize the role of immune responses. Microabscess formation in women with shorter lactation history brings to mind milk stasis as a contributing factor. As is illustrated in figure 1, milk interstitial extravasation and development of autoimmunity seem to form the key pathogenic pathway [21]. Hence, the authors recommend the term ‘milk-induced autoimmune’ to be used instead of ‘idiopathic’ (granulomatous lobular) mastitis.

Some authorities argue that in the case of reconstructive surgery, implants are best avoided as they themselves may act as foreign bodies and trigger further granulomatous inflammation [21] (fig. 2).

It should be noted that our pathologic reports were not made in a consistent and systematic fashion; hence, the relative frequency of the documented features (table 1) would be largely suggestive.

No uniform therapeutic protocol has been established for IGLM [10]. Surgical excision aiming at total removal was the traditional approach [22], but today the likely autoimmune nature [10] has made corticosteroids the mainstay initial treatment [23]. Oral steroids are effective and control the lesions in most cases. However, the disease is frequently recurrent and sometimes refractory, and steroid-sparing agents would be needed. Surgical resection is occasionally inevitable [24]. Generally, patients need a course of 3–6 months of systemic corticosteroids to achieve remission or cure [1]. An initial empiric antibiotic treatment is irresistible and was performed in 78% of our cases. We then put our patients on low-dose steroids (none of the patients required more than 35 mg/day to achieve a remission) and tapered treatment within 6 months. We finally achieved 90% remission or cure. Relapse and long-term steroid complications are the challenges of IGLM, and some have advocated methotrexate and azathioprine as maintenance modalities. We had a 16% relapse rate which is not unusual; relapse has been reported from as low as 8% to 23% and higher [25, 26]. The variation can be attributed to different follow-up and the frequency of resorting to surgical excision as the primary intervention. One additional cause for relapse might be that the disease is not completely in remission while steroids are being discontinued. Studies have shown that sonography has the best sensitivity in following the disappearance of the lesion (as compared with magnetic resonance imaging and clinical examination) [1]. We did not use methotrexate and azathioprine despite their usefulness as reported in the literature. Cytotoxic agents not only spare steroid complications but also reduce the number of indications for excision and the need for reinstitution of steroids [13, 24]. The authors acknowledge that a closer relationship should have been established with the oncological team.

Mastectomy is best avoided in IGLM [3] but still has its own indications for abscess and fistula formation, persistent wound infection, and refractory cases. 44% of our patients underwent surgery as the primary intervention, and 11.5% of the remaining eventually received surgical intervention following medical treatment (collectively, 55.5%). It should be noted that the classical approach of complete surgical excision (with or without steroids) [27] is now being replaced by primary medical therapy (steroids and immunosuppressive agents) [1] (our approach also evolved from primary surgery to primary medical treatment in the time course of the study).

In conclusion, IGLM should be included in the differential diagnosis of breast mass in patients of childbearing age, specifically in the years following pregnancy. A systemic survey needs to be carried out in order to best differentiate and diagnose the relevant infectious and coincident autoimmune entities. Diagnostic and therapeutic issues and the major differential diagnosis, i.e. malignancy, mandate a multidisciplinary approach covering radiology, rheumatology, oncology, gynecology, and surgery. We have drafted a preliminary clinical decision-making guideline for IGLM (table 2 and fig. 2).
Acknowledgment

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Disclosure Statement

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References