Gelatinous transformation of bone marrow: rare or underdiagnosed?

Khushdeep Kaur Shergill\textsuperscript{a}, Gagandeep Singh Shergill\textsuperscript{b}, Hari Janardanan Pillai\textsuperscript{c}

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\section*{ABSTRACT}

Gelatinous transformation of the bone marrow (GTBM) is a rare hematologic entity, which was first described by Paul Michael in 1930. GTBM is mostly associated with caloric intake/anorexia nervosa, although it also has been described accompanying other pathologic conditions, such as malignancy, systemic lupus erythematosus and HIV infections. Even though the diagnostic features of the hematopoietic tissue, such as hypoplasia, adipose cell atrophy, and deposition of a gelatinous substance in the bone marrow (which stains with Alcian blue at pH 2.5) are quite specific, the underlying pathogenic mechanisms remain poorly understood. Considering the evidence of reversibility—notably in cases of malnutrition and anorexia—this entity should be kept high on cards as a possible differential diagnosis of patients presenting with cytopenias and associated weight loss or starvation, especially in developing countries with nutritionally deprived populations. On an extensive review of the literature aimed at comprehensively addressing the evolution of the GTBM from the past century until now, we conclude that the lack of clinical suspicion and awareness regarding this pathologic entity has led to misdiagnosis and delayed diagnosis.

\textbf{Keywords:} Bone Marrow Disease; Bone Marrow Examination; Bone Marrow.

\section*{INTRODUCTION}

Gelatinous transformation of bone marrow (GTBM)—also known as starvation marrow—gelatinous degeneration and serous atrophy of bone marrow is a well-defined hematologic entity.\textsuperscript{1,2} Although the first references were noticed in works of Virchow (1821–1902), Herter (1865–1910), and Adami (1862–1926), it was Paul Michael, in 1930, who first reported 11 cases of GTBM in the bone marrow of autopsied cases.\textsuperscript{1} A large number of studies published at the end of the 20th century worldwide, considered GTBM as a histologic pattern originating from an associated cause rather than being a disease entity by itself.\textsuperscript{3}

In fact, GTBM has been described in association with a variety of clinical scenarios, chiefly in the young with malnourishment and anorexia.\textsuperscript{3,4} In cases with high clinical suspicion, the diagnosis can be easily established by studying the bone marrow aspirate or bone biopsy with Alcian blue staining at pH 2.5, which loses positivity after pre-treatment with hyaluronidase.\textsuperscript{3} The reported rare incidence of GTBM might just be the tip of the iceberg, with the lack of clinical suspicion being the most probable culprit for so few diagnoses.

This review was undertaken to study the epidemiological behavior and incidence of GTBM reports over the years while focusing on clinical
profiles, hematologic parameters and investigations, and proposed pathogenic mechanism.

MATERIAL AND METHODS

In our pre-specified protocol, we used the keywords “gelatinous transformation/degeneration of the bone marrow, starvation marrow, and serous atrophy of the marrow” to identify the target papers for a systematic literature search. The databases, PubMed, Medline, Google Scholar and Research Gate, were accessed to retrieve the articles published until June 2017.

Case and series reports, original research articles, images, letters to editors with titles having the aforementioned keywords were included in the study. Data including the first author’s last name, publication year, country of origin and total number of cases reported in each study was compiled. All open access articles and abstracts of subscribed articles were included in the study.

Articles published in a language other than English for which no translation or English abstract was available, were excluded.

All articles fulfilling inclusion criteria were reviewed in duplicate and independently, and data extraction was held by two authors (KKS & GSS).

All articles were assessed for following variables: total number of studies published with total number of cases; clinical profile including age, sex and the associated disease; altered hematological indices; diagnostic work-up modality (cytological and/or histologic parameters of bone marrow); reversal or improvement with treatment; and additional information reported with respect to the pathogenesis of GTBM by the different authors.

Data were compiled in chronological order (decade wise); however, no quantitative data synthesis or statistical analysis could be performed due to considerable clinical and mathematical heterogeneity of the compiled data.

RESULTS

Our literature search gathered 79 papers (N) with relevant titles, amounting the total of 522 reported cases (n) as shown in Table 1. Of these 79 papers, 66 were in the English language while 13 were in other languages, such as French, German, Dutch, and Polish. Of these 13 studies, 4 had an English language abstract. Hence the remaining 09 articles (n = 26) were excluded.

A final list of 70 titles was compiled, which comprised 48 full articles (n = 394) and only 22 Abstracts (n = 102). Out of these, 48 open access articles were subjected to review by two authors independently and the remaining 22 subscribed articles only with abstracts were reviewed for available data.

We observed that the research articles on GTBM, which constituted the bulk of cases, had limited information on the parameters of the individual cases. Hence, the scope for statistical analysis of data was limited. However, an overall trend was established based on the available data.

Incidence of Gelatinous Transformation of Bone Marrow Over the Last Century

The first available published study on GTBM by Paul Michael (1930) comprised a total of 11 cases. Over the following three decades, no published data were available. In 1967, a study by Pearson surfaced, reporting GTBM in three cases of anorexia nervosa. In the subsequent decade; two more studies were published by Tavassoli et al. (1976) and Seamen et al. (1978) who described the condition in 9 and 14 cases, respectively. In the 1980s, six more studies were published, describing the condition in seven different patients. However, no open access article could be found on a literature search of the archives till 1990. The number of studies tripled (N = 18) in the 1990s with a total of 52 cases reported, including one original research article by Mehta et al. who described the condition in 22 patients with HIV—the first ever study from India—by Ray et al. in 1992, and another one by Basu et al. At the dawn of the 21st century, a deluge of research articles on GTBM were published. Among them, 21 new studies appeared in the first decade itself, including the largest study by Bohm who studied 80,000 bone marrow samples and reported the GTBM incidence of 0.2%. In the last 7 years, 30 new studies comprising 117 cases were published. Thus, a total of 79 studies with 522 individual cases of GTBM have been published in the last 87 years (Table 1). These data clearly indicate...
Table 1. Data sheet of cases with GTBM (1930-2017)

| Year          | Number of studies (n) | Number of cases (n) | Age groups (<15: 15–40: 40–60: >60) | M:F ratio | Associated entity, number of cases | Hematologic characteristics | Diagnostic work-up | Follow-up |
|---------------|-----------------------|---------------------|-------------------------------------|-----------|-----------------------------------|----------------------------|---------------------|-----------|
| 1930-1939     | 1                     | 1                   | (Average age: 40 years)             | 10:1      | Tuberculosis: 4, Malignancy: 3, Miscellaneous: 4 | Anemia: 10/11, Leukopenia: 1/11 | BMB                  | -         |
| 1940-1959     |                       |                     | Nothing published                   |           |                                    |                            |                     |           |
| 1960-1969     | 1                     | 3                   | 2:1                                 | 0:3       | Anorexia Nervosa: 3                | 1 case: leukopenia          | BMA: 2 cases; BMA & BMB: 1 case | Response to treatment |
| sa1970-1979   | 2                     | 23                  | 0:1:6:10                           | 11:6      | Malignancy: 7 Anorexia/ Malnutrition: 6 Miscellaneous: 4 | At least 1 cytopenia; mostly anemia | BMA & BMB             | Improved with nutrition in animals |
| 1980-1989     | 6                     | 7                   | NA                                 |           | Anorexia: 5 Malignancy: 1 SLE (overlap): 3 | At least 1 cytopenia; Anemia most common, Pancytopenia: 1 case | -                    | -         |
| 1990-1999     | 18                    | 52                  | Complete data                       | NA        | Anorexia/ Malnutrition: 10 HIV: 22 | Mostly anemia              | Post-chemo therapy: 10 |           |
| 2000-2009     | 21                    | 309                 | 50:106:36:83                       | 150:110   | Anorexia/ Malnutrition: 79 Malignancy: 66 Infections: 53 Aplastic anemia: 17 Miscellaneous: 47 | 2 cases without cytopenia Mostly anemia Pancytopenia: 2 | BMB & BMA: 70 cases BMA: 54 cases BMB: 179 cases | 23/36 cases improved (5 with nutrition, 1 with EPO) |
| 2010-2017     | 30                    | 117                 | 18:3512:10 (30 cases btw 26 & 66 yrs) | 65:40     | Anorexia/ Malnutrition: 44 Malignancy: 22 HIV: 9 Infections: 7+/–15 | 2 cases without cytopenia Most cases with anemia, Pancytopenia: 14 | BMA: 774 BMB: the remaining |           |
| TOTAL         | 79                    | 522                 | 70:143:54:103                      | 236:160   | Anorexia/ Malnutrition: 147 Malignancy: 109 | -                          | -                    | -         |

BMA = bone marrow aspirate; BMB = bone marrow biopsy; btw = between; EPO = erythropoietin; GTBM = gelatinous transformation of bone marrow; NA = not available; PRBCs = packed red blood cells; SLE = systemic lupus erythematosus. *No full published article was available during this time period; hence, data interpretation is limited. **Mehta et al. reported in 22/75 cases of HIV (86% males, mean age: 33 years; all were anemic, 50% cases hypocellular, aspirate more difficult). ***Jain et al. (India) studied 43 cases, 14 of which were children; mostly presented anemia and malnourishment.
a rising trend in reporting the condition, especially in the last two decades.

Clinical Profiles

Our analysis showed a total of 236 male patients compared to 160 females with a M:F ratio of 1.5:1. Cases were divided into four age groups: <15 years, 16-40 years, 41-60 years and >60 years. In the initial studies, the most commonly involved age group was 40-60 years and >60 years with an occasional case in the <15 years age group. However, with more cases of young patients being reported in the past two decades, a total of 143 cases were noted in the age group of 15-40 years, followed by the elderly age group of >60 years (103).

In our analysis, a total of 70 cases were seen in the <15 years age group (Table 1). The most common reported etiologies associated with GTBM were anorexia nervosa, and malnutrition or malabsorption with weight loss (47 cases). Similarly, the majority of cases, reported by Jain et al.4 (the largest case series from India), also concluded that malnutrition is the most common condition associated with GTBM. Further GTBM associated with malignancy was reported in 109 cases and comprised several types of cancers, such as hepatocellular carcinoma,64 oral cancers,74 along with primary malignancies of the bone marrow, such as acute myeloid leukemia (AML),19 and chronic myeloid leukemia (CML)47. Another 60 cases were associated with different types of infective agents excluding those reported in HIV patients.16 A total of 17 cases of aplastic anemia with GTBM were reported in study by Sen et al.37 As in cases of malignancy, HIV infection and chronic infections, like tuberculosis, where weight loss is quite often an associated complaint, there seems to be an overlap leading to an underestimation of cases of GTBM associated with anorexia and malnutrition. A total of 40 cases were labeled as miscellaneous, which comprised different associated diseases, such as systemic lupus erythematosus,15,49 visceral Leishmaniasis,17 hemochromatosis,48 and metabolic abnormalities, such as hyperthyroidism, adrenal insufficiency,74 pyrexia of unknown origin, and alcoholic pancreatitis have also been published.

Investigations

The most common altered laboratory parameter found in the majority of the cases was the presence of at least one peripheral cytopenia. In this setting, anemia was the most common. In some of the larger series, almost 100% of the cases were reported to have anaemia.3,16,37,63 Pancytopenia as the initial presentation was present in 22 cases.58,64,65,68,70,73 However, cytopenias were occasionally lacking,41,71 and a single case of myelofibrosis, accompanied by GTBM, had leukocytosis along with thrombocytosis.31

In all cases, the diagnosis was initially achieved either on a bone marrow aspirate or bone marrow biopsy; however, the bone marrow biopsy was reported to be the method of choice mainly in cases with a dry tap on bone marrow aspiration.3,63 Histological examination was characterized by different patterns of bone marrow cellularity, varying from hypocellularity to normocellularity—or even cases of hypercellularity with focal areas of hypocellularity surrounding the gelatinous deposits.31,63,65,66 Böhm3 graded the bone marrows from 1 to 4 based on the severity of gelatinous transformation.

Bone marrows were stained with Alcian blue at pH 2.5 along with routine H&E stains and additional stains like Congo Red, and periodic acid–Schiff to rule out other differential diagnoses.

Prognosis

Pearson5 and Tavassoli et al.6 firstly reported the reversibility of GTBM on animal experiments. In our review, we found 35 cases that showed the disappearance of the GTBM on follow-up on repeated biopsy/aspirate after the initiation of treatment, which was either in the form of nutritional diet (15 cases),58,60,62,66 packed red blood cell (PRBCs) transfusion (one case),57 granulocyte colony stimulating factor (G-CSFs) (three cases)51 or treatment of the underlying disease.

DISCUSSION

Since 1930, there has been an increasing number of reported cases of GTBM. From the advent of the 21st century, more than 50 studies were published accounting for more than 400 new cases (Table 1). The majority of these cases were from countries like India where GTBM was unheard of before the 1990s. The reported incidence of GTBM varies from 0.2%,3 to 4.4%37 and 4.8%.66 This rising trend in its incidence over the years indicates that GTBM, in the
current scenario, is anything but rare. On review of the literature, Seaman et al.\textsuperscript{2} seems to aptly describe GTBM as “Not uncommon,” whereas the majority of studies continue to report it as “a rare condition.” Based on our analysis, we conclude that the low incidence observed in the 20th century can be ascribed to the lack of awareness about the existence of such an entity.

Overall, there is a male preponderance in the cases reported so far.\textsuperscript{3,37} However, in certain studies, like Abella et al.,\textsuperscript{36} all 22 cases reported were females while in study by Mehta et al.,\textsuperscript{16} 86% cases were males. This may probably be due to a selection bias in such studies. Young adults and the elderly constitute the most commonly affected age;\textsuperscript{3} however, there has been an increase in the number of cases reported in the pediatric age group over last two decades, most of which have been reported from India.\textsuperscript{4,37,65} Hence, it is noteworthy that no age group is immune to the condition, and thus GTBM should not be neglected as a differential diagnosis in any age group.

The underlying disease spectrum associated with GTBM is heterogeneous. The most common association had been reported with cases of malnourishment/starvation resulting from various causes like anorexia, chronic infections, or malignancy.\textsuperscript{1} Though most of the initial case reports were associated with non-hematopoietic malignancies, recently GTBM has been reported in association with various hematopoietic malignancies like AML,\textsuperscript{14,18,25} acute promyelocytic leukemia (APML) on All Trans Retinoic Acid,\textsuperscript{70} acute lymphocytic leukemia (with and without dasatinib),\textsuperscript{61,63} CML on imatinib,\textsuperscript{47,55,56,72} CML in chronic phase,\textsuperscript{50} myelodysplastic syndrome,\textsuperscript{45,59} multiple myeloma\textsuperscript{33} and myelofibrosis,\textsuperscript{31} which also have been published from all over the world. In many of these cases, marked weight loss with anorexia was present,\textsuperscript{44,63,74,77} thus pointing to the possibility that the underlying mechanism resulting in the development of GTBM remains the same as in the cases of chronic infections and malignancies, anorexia, and malnutrition.

A remarkable number of patients had bi- or pancytopenia at the initial presentation, and almost all had anemia, irrespective of the underlying associated disease. Even though no correlation between the severity of GTBM and the peripheral hematological parameters have been reported,\textsuperscript{3,63} it still should be kept in the differential diagnosis of cytopenias. Regarding the investigation modality, bone marrow biopsy appears to be the gold standard, and it appears preferable to perform trephine biopsy along with aspirate in suspected cases rather than doing aspirate alone. In 2010, Sims\textsuperscript{80} described the characteristic findings of patchy areas of hyperintensity in T2-weighted images in cases of GTBM detected by magnetic resonance imaging (MRI). However, there are cases of GTBM misdiagnosed as spinal tumours\textsuperscript{12} or misinterpreted as technical errors on MRI.\textsuperscript{75}

The diagnosis is easily achieved in bone marrow biopsy, which exhibits the features of hematopoietic hypoplasia, adipose cell atrophy, and the deposition of gelatinous substance in bone marrow that stains with Alcian blue at pH 2.5 (Figure 1).\textsuperscript{1,3,6} However, a large number of cases present normocellular or hypercellular marrows or focal areas of hypocellularity surrounding the gelatinous substance, which easily can be misinterpreted as bone marrow edema, necrosis, or amyloid. The misinterpretation occurs with a lack of suspicion or knowledge of this entity. (Table 2).\textsuperscript{3,69} Recently, GTBM post-chemotherapy has been reported without fat atrophy.\textsuperscript{18,33} Although the pathogenesis of GTBM remains unknown, the majority of the studies, especially in cases of nutritional deprivation, agree upon a common pathophysiological mechanism of fat mobilization in the face of starvation followed by the deposition of hyaluronic acid. This excessive accumulation of extracellular gelatinous material interferes with the hematopoietic microenvironment leading to the suppression of hematopoiesis, which is reflected as peripheral blood cytopenias.\textsuperscript{51,59}

In GTBM associated with infectious diseases, as in HIV infection, the secretion of cytokines, such as interleukin (IL)-1, IL-2, and the tumor necrosis factor, are considered to be the driving factors.\textsuperscript{43} Similarly, it may hold true for the cases associated with malignancy. Alternate theory suggests that the tumor cells may play a role in the initiation of hyaluronic acid formation.\textsuperscript{11} In the cases of GTBM following chemotherapy, the inhibition of tyrosine kinase activity leads to the blockage of downstream signal pathways affecting extracellular matrix deposition, and adipocyte differentiation has been proposed as the possible mechanism.\textsuperscript{72} Local milieu factors of the marrow as the systemic “stress factors” have been proposed as the inciting factors for the development of GTBM in cases associated with aplastic anaemia.\textsuperscript{37}
Long-term outcomes and the prognosis of GTBM have not been well documented yet. A few authors have stated that GTBM, by itself, does not have any prognostic significance, and that the survival of patients who develop GTBM is dependent on the nature and stage of the underlying disease at the time of the diagnosis. However, with the increasing number of case reports showing an improvement of the GTBM in patients after balanced nutrition, PRBCs, G-CSFs, erythropoietin, the irreversibility of GTBM has been questioned. Mant and Faragher reported that GTBM is a temporary condition, which reverses on balanced nutrition in cases of anorexia nervosa. The timely diagnosis and early treatment in such cases would reduce the costs of long-term health care expenses, especially in developing countries, like India.

Table 2. Differential diagnosis of GTBM

| Bone Marrow | GTBM | Marrow Necrosis | Marrow Edema | Amyloid | Aplastic Anemia |
|-------------|------|----------------|--------------|---------|----------------|
| Microscopy  | Extracellular eosinophilic amorphous material with fibrinous precipitate. Inverse relationship with fat vacuoles. Vacuole outlines prominent, ragged and irregular. Diffuse hypoplasia to only focal hypoplasia with surrounding areas showing normal cellularity | Eosinophilic granular material with the presence of cellular karyorrhectic debris | Hypocellular marrow areas with fat cells of normal size and in normal quantity | Homogenous pink material in the vessel wall, or interstitial with intact fat vacuoles. There is an increase in plasma cells | Loss of myeloid tissue without any fat cell atrophy |
| Congo Red   | -    | -              | -            | +++     |                |
| Alcian blue pH 2.5 | +++ (Stains blue) | -              | -            | -       | -              |
| PAS         | Stains pink | -              | Pale pink    | -       |                |

GTBM = gelatinous transformation of bone marrow; PAS = periodic acid-Schiff.
CONCLUSION

From an extensive review of the literature, we conclude that GTBM is a condition that is underdiagnosed because of the lack of clinical suspicion. Though large-scale prospective studies are required to establish the pathogenesis, GTBM should be considered as a differential diagnosis in all cases presenting with weight loss, malnutrition, and refractory peripheral cytopenias—irrespective of age profile and associated co-morbidities.

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**Correspondence**
Khushdeep Kaur Shergill
Department of Pathology, Armed Forces Medical College (AFMC)
Pune - Maharashtra - India - 411040
Phone: 070303991112
shergillkhushdeep@gmail.com