Numb Chin Syndrome: Diagnostic Significance Over 57 Years and Review of 550 Patients in the Literature Worldwide

Keywords: Numb chin syndrome; Mental neuropathy; Mental nerve numbness; Lymphoma; Leukemia; Brain; Bone; Breast; Prostate; Cancer

Abstract

Numb Chin Syndrome (NCS) is a heterogeneous condition of mental nerve neuropathy, playing in concert with multiple disease mechanisms, but all result in a disruption of unilateral or bilateral facial nerves, fever, and swelling of the lower lip. The NCS has been reported as a prodromal symptom of pending malignancies, a possible precursor of cancer or leukemia, and sometimes merely a benign, painful or annoying condition. However, any causal relationships between the NCS and malignancy have not been made, nor aggregated, and therefore remain unclear. This thorough review of all the available current literature in the PubMed interface was performed by using the key words, “mental neuropathy,” “mental nerve numbness,” and “n umb chin syndrome.” There were 186 peer-reviewed articles reporting 550 patients from 28 countries worldwide including 150 benign, 398 malignant, and 2 unknown cases. The seven most common malignancies from high to low frequency were: lymphoma, bone cancer, oral cancer, breast cancer, brain cancer, leukemia, and prostate cancer. While the clinical implications of the NCS are not fully understood, this delineative study will aid physicians confronted and confounded with various malignancies, neuropathies, and cytogenetic abnormalities of patients initially presenting with the symptom of a numb chin.

Abbreviations

HIV: Human Immunodeficiency Virus; MRI: Magnetic Resonance Imaging; NCS: Numb Chin Syndrome; USA: United States of America

Introduction

Numb Chin Syndrome (NCS) is a set of neuropathological symptoms including lower facial numbness, due to paralysis of the distal trigeminal nerve, especially right, unilateral, dysesthesia or paresthesia (numbness) of the chin, local fever, swelling, redness, persistent pain, a thickening sensation of the lower lip, and dullness in the mandible. When contemplating this worldwide, not too rare disease of the numb chin, a mental image of Rodin’s famous bronze statue, “The Thinker” comes to mind. He is pressing the back of his right hand to the right side of his chin, right elbow to his left knee. Is he applying pressure to alleviate some pain, feeling numbness in his chin, contemplating some kind of worsening condition? Or what? We wonder. Sitting in this position, what is he thinking?

In the beginning in London, Charles Bell first described this syndrome in 1830 in a woman with breast cancer and noting mental nerve anesthesia in patients presenting with trauma to the jaw and retromandibular tumors [1]. In the United States of America, numb chin was first mentioned as a syndrome, in 1963 as an innocuous symptom, albeit an ominous one, that would indicate metastasis and manifest malignancy [2]. In Japan, the NCS was first reported in 1989 as the initial manifestation of a case of Burkitt’s lymphoma [3].

Now, 190 years after Bell’s first mention of it [1], we have attempted to determine if the fulminating numb chin is an ominous sign of an upcoming malignancy, a prodromal syndrome of a life threatening disease, or merely a benign but painful nuisance. Although there have been various reviews reporting malignancies and diseases associated with a numb chin, they focus on certain kinds of malignancies, such as lymphoma, leukemia, lung, breast, brain, bone, and prostate cancers, among other major noncancerous maladies. The aim of this study was, therefore, to summarize all of the reported cases of the NCS from all over the world for the last 57 years, looking into benign and malignant cases, to determine if there were any patterns that may suggest or even prove causality.

Subjects and Methods

An NCS review was performed of all the available current literature in the PubMed interface, during the 57-year period from 1963 to 2020. The key words “mental neuropathy,” “mental nerve numbness,” and “n umb chin syndrome” generated a considerable body of literature comprised of 186 relevant, peer-reviewed articles. The reported cases were manually counted for the various signs and symptoms, histories, diagnoses, treatments, prognoses, and actual outcomes.

Results

A retrospective analysis of 186 articles revealing 550 patients, who presented with signs and symptoms of the NCS at 122 sites in 28 countries throughout Europe (n = 198), North and South America (n = 177), the Middle East (n = 107) and the Asia/Pacific region (n = 68), composed this exhaustive study. The range in patients’ ages was from
were HIV positive. Noteworthy, there were 11 cases of NCS related to or Burkitt’s leukemia; and 3 of those 13 Burkitt’s leukemia patients were diagnosed as having Burkitt (non-Hodgkin) lymphoma and/or Burkitt’s leukemia. Moreover, 2 children, 12 years old and 18 years old respectively, were reported as having succumbed to their illnesses in the most recent published report. In Japan, the oldest and youngest patients presenting with the initial symptoms of the NCS were a 73-year-old patient with prostate cancer and an 18-year-old boy with Burkitt’s lymphoma/leukemia. They presented with 2 weeks of bilateral lower limb pain and left chin numbness. She was diagnosed with the NCS secondary to mandibular cell arteritis, also called temporal arteritis, successfully treated with oral Prednisolone [4]. There was only 1 case reported of recurrence, and that being one of multiple recurrences, four fulminant painful but benign attacks, over a period of 16 years [5]. Currently, at 70 years of age, the 5-year follow-up revealed that patient to be asymptomatic and in good health.

Discussion

This study showed that a numb chin and a fat lip correlated with malignancy more often than might otherwise be expected in as many as 398 (72%) of 550 NCS patients. First mention of the syndrome of the numb chin in the United States of America was in 1963 [2]. In Spain, in 2008, it was reported that this symptom may be the first manifestation of systemic cancer [6], a symptom of spread of an established tumor, or a sign of infiltration in an intraoral lesion [6]. Moreover, in that study, 18 of 22 patients (82%) died before a mean of 20 months after the initial presentation, further showing that these symptoms are mostly indicative of very poor prognoses [6]. By 2014, only 6 years later, and as many as 83 cases had been reported worldwide; and reported in Germany it was reported NCS was a “rare” and often overlooked symptom of extracranial malignancies [7].

Beginning in 1963 through 2018, 546 cases were reported around the world in those 55 years, amounting to an average of 10 new NCS cases being reported annually. Notwithstanding, Malaysia and Japan reported the syndrome as seemingly rare in their countries but manifesting malignancy and morbidity [8,9]. In Malaysia, following a mastectomy for stage 2 breast carcinoma, a 51-year-old woman presented with 2 weeks of bilateral lower limb pain and left chin numbness. She was diagnosed with the NCS secondary to mandibular tumour metastasis, started on palliative chemotherapy and radiation, and eventually discharged with the only lingering complaint of a numb chin [8]. In Japan, the oldest and youngest patients presenting with NCS symptoms were a 73-year-old patient with prostate cancer and an 18-year-old boy with Burkitt’s lymphoma/leukemia. They both were reported as having succumbed to their illnesses in the hospital [9].

Collectively these results, from 28 countries in this 57-year longitudinal worldwide study of 186 articles in the medical literature reporting NCS events and outcomes, could be important to physicians with respect to performing further examinations to make more accurate diagnoses and earlier treatment decisions for patients presenting with the initial symptoms of the NCS. The study revealed that patients presenting with a numb chin, intractable pain and a swelling, redness, persistent pain, a thickening sensation of the lower lip, ought to be carefully followed...
up for fear of future maladies. Physicians in Germany, Japan, and Italy, asked if this syndrome was a reflection of malignancy, an immune-mediated disease, or a harbinger of medication-related (e.g., bisphosphonate), osteonecrosis of the jaw and methotrexate-associated lymphoproliferative disorders [7, 10, 11].

Even though the mechanism of this neurological phenomenon of the numb chin remains unclear to date, and therefore often controversial, a significant majority of cases have led to the comorbid prognoses of malignancies, as evidenced in these 398 patients including 63 (11%) morbidities. In terms of perspective only, the 11% rate of death for NCS patients is significantly higher than the worldwide

| Cancer types         | n  | %  |
|----------------------|----|----|
| Lymphoma             | 61 | 14 |
| Bone                 | 55 | 13 |
| Oral                 | 51 | 12 |
| Breast               | 40 | 9  |
| Brain                | 26 | 6  |
| Leukemia             | 25 | 6  |
| Prostate             | 20 | 5  |
| Adenocarcinoma       | 16 | 4  |
| Lung                 | 12 | 3  |
| Multiple myeloma     | 11 | 3  |
| Sarcoma              | 4  | 1  |
| Skin                 | 2  | 0.5 |
| Colon                | 3  | 1  |
| Bladder              | 2  | 0.5 |
| Esophageal           | 2  | 0.5 |
| Stomach              | 1  | 0.2 |
| Liver                | 1  | 0.2 |
| Adenoid              | 1  | 0.2 |
| Renal                | 1  | 0.2 |
| Adrenal              | 1  | 0.2 |
| Uterine              | 1  | 0.2 |
| Mediastinal          | 1  | 0.2 |
| Plasmocytoma         | 1  | 0.2 |

| Cancer type unknown  | 85 | 20%|
|----------------------|----|----|
| Total                | 425| 100|
| Multiple malignancies| 37 | 9  |
| Multiple malignancies| 37 | 9  |

*Reference numbers are not from the text but from the Appendix (available upon request).
Each reference number represents one case unless noted otherwise in parentheses. Unknown (i.e., not reported)

With hundreds of malignancies the world over, the NCS signs and symptoms were revealed to be prodromal, the first manifestations in many other severe and debilitating diseases. Of those, perhaps the most debilitating is multiple sclerosis [7, 12–15]. Fortunately, one case of multiple sclerosis presenting with the symptoms of the NCS in Germany was successfully treated with interferon-B [14]. In France, a successful treatment with rituximab was reported in a patient with mental nerve neuropathy in primary Sjogren’s syndrome, the autoimmune disease often comorbid with other immune diseases, especially rheumatoid arthritis and lupus [16]. In the USA, the NCS was the initial presentation of a posttransplant lymphoproliferative disorder [17]. In Italy, there were 3 cases of sudden violent, diffuse pain in the premolar region reported [18]. In the USA, a patient successfully underwent a transoral surgical resection of an elongated, calcified, styloid process to relieve the mental nerve paresthesia and pain caused by the dynamic compression of the alveolar inferior nerve [19]. In Spain, the NCS was secondary to a solitary schwannoma [20]. And in Japan, a 39-year-old woman suffered a large solitary fibrous tumor on the infratemporal fossa [21]. Thus, as is well evidenced, all over the world, there have been and still are hundreds of cases of NCS as the initial, primary sign of oncoming maladies, the degree of severity of which becomes evident in time. Therefore, early diagnoses and adequate treatment are often the difference between life or, if neglected and undiagnosed, a possible lingering death to cancer – the difference between remission-free survival or morbidity.

Burkitt lymphoma/leukemia is the most frequent subtype of hematological malignancies in the NCS. It is highly aggressive and often presents in extranodal sites and/or as acute lymphoblastic leukemia. A leukemic phase can be observed in patients with bulky disease, typically in males, and clearly present as leukemia with peripheral blood and bone marrow involvement. In the 2016 revision of the fourth edition of the World Health Organization
classification of myeloid neoplasms and acute leukemia, Burkitt leukemia was classified as a subtype of Burkitt lymphoma [22]. Three epidemiological subtypes are recognized, which mainly differ in geographic distribution. Endemic Burkitt lymphoma occurs in equatorial Africa and Papua New Guinea, where the jaw and other facial bones are typically the sites of presentation in approximately 50%-70% of those cases [23,24]. While not reported per se, it is conceivable that NCS signs and symptoms are physical findings in endemic Burkitt lymphoma patients. However, sporadic Burkitt lymphoma and immunodeficiency-associated Burkitt lymphoma tumours in facial structures, particularly in the jaw are rare. Furthermore, lytic destruction of an area of bone due to myeloma and lymphoma is a common symptom because leukaemia and multiple myeloma proliferates in the bone marrow. This explains the physiological causality and neuropathic association that the NCS is often observed in patients with various types of hematological malignancies.

In these 57 years, there were 29 reported cases of the numb chin suspected of being an outright prodromal symptom of various malignancies. However, as an example of the reverse occurring, a case was reported of a 22-year-old man who after achieving a complete 5+year remission of CD+20 positive precursor B-ALL, presented with gradually progressive altered mentation, slurring of speech, and chin paresthesias over a 72-hour period [25]. His physical exam revealed poor concentration, dysarthria, diminished sensation over the mental area, and sialorrhea; however, the remainder of the examination was unremarkable. There was another case, reported in 2015, of NCS being the secondary symptom to cancer that had started from gastric adenocarcinoma and metastasized to the brain as leptomeningeal carcinomatosis [26]. All the other 548 cases in the present series were of NCS being the first forewarning, an alarming-type symptom of what usually turned out to be a malignant and/or metastasizing disease. Sometimes for physicians unfamiliar with the NCS, these symptoms were apparently not taken as a red flag or did not sound an alarm and malignancies or debilitating diseases were manifested at later stages in the patients’ prognoses.

Remarkably, of 550 cases, 150 (27%) were benign. Moreover, in the course of compiling these data, it was conjectured that a significant number of NCS cases worldwide must go unreported and/or lost to follow-up. There were 10 articles that did not include any numbers of patients but cautioned physicians about the various aspects of the NCS and advised them to be on the lookout for these symptoms. After a 55 year period with an average of 10 new NCS cases reported annually, the reporting of NCS cases seems to have gone silent in 2019 because, to our knowledge, there were no new cases reported until only one in 2020. That case was in Qatar, of a 30-year-old Indian man with acute myeloid leukemia who, after being treated and discharged, returned to his home country and was lost to follow-up [27]. The facts and outcomes of these 550 cases from 28 countries on all 6 continents from around the world, corroborate those in 2014, in Germany, when it was reported that both benign and malignant diseases cause the symptoms of this syndrome to flair up, that there are many misdiagnoses, and, in a large majority of patients, the symptoms often lead to a revelation of malignancy [7].

When a patient presents complaining of a numb chin, and/or dyesthesia, fever, and a swelling sensation of the lower lip, the overwhelming findings of this report will remind physicians to see this set of signs and symptoms as “a red flag” and follow up with sufficient examinations allowing them to make the right call in the early stages of diseases to help mitigate pending threats. Therefore, we hope that this study helps guide physicians to delve further into the reaches of the human nervous system to elucidate the relationships among the NCS, paralysis of the distal trigeminal nerve, leukaemia, lymphoma, bone, oral, brain, breast, and prostate cancer, metastatic adenocarcinomas, and various other confounding, life-threatening neuropathies and cytogenetic abnormalities.

Study limitations

For the purposes of compiling these data and “bean counting” in this report, we have attempted to extrapolate any and all duplicated cases in the literature. Likewise, in some articles, there were omissions discovered wherein the exact data was not reported. However imperfect, therefore, further quantitative and qualitative considerations, which may logically and scientifically apply to these data, can be made by each individual physician.

Conclusion

This 57-year, 28-country, worldwide study of 186 articles revealed 398 (72%) malignancies in 550 patients who suffered the fulminant, fullblown symptoms of the NCS. Because significant evidence of causality remains elusive, this research opens up avenues for future study, specifically, to examine how the NCS status affects the prediction of numerous and seemingly unrelated maladies and malignancies. With a closer look at this syndrome, early detection may reduce long hospital stays for patients suffering with cancer, and some deaths may be avoided.

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Acknowledgements

We thank Michiel Brandt, MA, Monterey Institute of International Studies, albeit posthumously, for her 30 years of being “high on life” and her dedication and inspiration that made this study possible. Her death was due to an 8-year bout with leukemia. We are also greatly indebted to Takashi Katakura, PhD, of the Department of Physiology, Kitasato University School of Medicine, for his technical assistance.