Abstract

Aggressive periodontitis is a low-prevalence, multifactorial disease, of rapid progression and with no systemic compromise. It presents immunological alterations, a strong genetic influence, familial aggregation and early onset. It can be localized or generalized. It is not clear whether it is an independent periodontal disease, or if it is the phenotypic expression of chronic periodontitis in susceptible patients. Its diagnostic protocol includes a dental medical history, a clinical periodontal examination and a radiological examination. Treatment usually includes improving oral hygiene, dental scaling and root planing, as well as systemic and local antibiotic therapy. Surgical therapy will depend on each individual case. Maintenance therapy is essential to achieve better results. The aim of this paper is to review diagnostic and therapeutic protocols, and to propose a treatment flowchart based on the latest scientific evidence.

Keywords: aggressive periodontitis, diagnosis, therapy, antibacterial agents, microbiology.

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INTRODUCTION

Aggressive periodontitis is a rare form of periodontal disease, which is characterized by rapid attachment loss, bone destruction, non-contributory medical history and family history of the cases\(^1,2\).

Early identification of this pathology can help prevent early loss of teeth. It is important to ensure appropriate care, prevention, diagnosis and treatment of aggressive periodontitis in health facilities in order to reduce the risk of functional impairment of the stomatognathic system and its systemic impact. It is also a priority to standardize and articulate the procedures provided in all the national healthcare institutions for this pathology.

This paper aims to be an updated review of the literature on diagnostic methods and treatments for aggressive periodontitis according to scientific evidence, which is expected to be useful for general practice dentists and periodontists.

METHOD

A non-systematic narrative review was conducted regarding the diagnosis and treatment criteria for aggressive periodontitis. The descriptors used were: “aggressive periodontitis”, “diagnosis”, “treatment”, “antibiotics”, “Microbiology”. Papers in English and Spanish were included. The papers were selected from PubMed, SciELO, Lilacs and Google scholar.

RESULTS

1. Definition. Aggressive periodontitis is a type of periodontal disease with rapid insertion and alveolar bone loss, which is characterized by familial aggregation and affecting healthy individuals, except periodontitis\(^3\). It usually affects young people, but it can appear at any age, although this is less frequent\(^3,4\).

2. Terminology and classification In 1999, the American Academy of Periodontology (AAP) Workshop coined the term “aggressive periodontitis”, and the disease was classified as “localized and generalized”\(^3,4\).

3. Etiology Aggressive periodontitis is a multifactorial and genetically complex disease. An increase in host susceptibility may be caused by the combined effect of genetic
predisposition, environmental factors (virulent pathogens, tobacco smoking, personal and professional hygiene) and local contributing factors. Furthermore, herpesvirus could be an additional factor of susceptibility and severity in aggressive periodontitis\(^9\).

The generalized form of aggressive periodontitis has been strongly associated with bacteria such as *Porphyromonas gingivalis*, *Agregatibacter actinomycetemcomitans* (Aa) and *Tannerella forsythia*\(^6,7\); recent studies suggest that other microbial species could also be associated\(^8\). In generalized aggressive periodontitis there is an inadequate response to the action of the periodontal pathogenic bacteria caused by a variety of genetic and immunological risk factors\(^9,10\).

Recent findings suggest that the pathogenesis of localized aggressive periodontitis is associated with severe abnormalities in the neutrophil function, producing neutrophil-mediated tissue injury\(^11\). These abnormalities appear to be the result of a stage of chronic hyperactivity of neutrophils; these findings are consistent with recent reports of decreased neutrophil chemotaxis in localized aggressive periodontitis\(^12,13\). Some studies suggest that this is due to abnormalities in the transduction signals. In addition, the neutrophils of patients with localized aggressive periodontitis show reduced calcium entry\(^14\), defective calcium influx factor\(^15\), abnormal activity of protein kinase C\(^16\), among other abnormalities.

4. Epidemiology

The prevalence of aggressive periodontitis is variable. It is estimated that there is a low prevalence (less than 1%) in Caucasian subjects living in developed countries compared to those living in developing countries (-0.5 to 5\%)\(^17\). One aspect that complicates the interpretation of aggressive periodontitis regarding epidemiological information is the parameters used to evaluate the identification of cases, which vary from one study to the other\(^18,19\). Africans and African-Americans seem to have the highest prevalence of aggressive periodontitis: 1.0-3.0\%. They are followed by Asians, with 0.4-1.0\%, and Hispanics and South Americans, with 0.5-1.0\%, compared to Caucasian young populations, with 0.1-0.2\%\(^20-22\). Löe and Brown estimated that African American adolescents are fifteen
times more likely to have aggressive periodontitis than white adolescents\textsuperscript{(23)}. Kissa et al. found no differences in prevalence by gender and ethnic group\textsuperscript{(24)}. Mwokorie and Arowojolu found a prevalence of 1.6\% in a population group whose age ranged from 17 to 34 in a Nigerian hospital\textsuperscript{(25)}. Albandar et al. reported high prevalence levels of aggressive periodontitis among Ugandan students aged 12-25; of which 6.5\% showed generalized or localized aggressive periodontitis; and 22\% showed incidental aggressive lesions\textsuperscript{(26)}. Hodge et al. suggested that genetic factors are more significant than history of smoking in the manifestation of generalized aggressive periodontitis\textsuperscript{(27)}. There are no epidemiological studies related to aggressive periodontitis in Perú.

5. Risk factors

One of the main risk factors of aggressive periodontitis is family history associated with inherited genetic traits. There is strong evidence that shows family history in young patients with early onset of aggressive periodontal disease\textsuperscript{(1)}. Efforts made in the last twenty years to identify specific genetic variations involved in the disease have not been conclusive.

Regarding oral hygiene, studies have suggested that there would be no correlation between plaque levels and the presence of disease\textsuperscript{(1)}.

Risk factors identified for periodontal diseases are similar to the ones for chronic periodontitis and aggressive periodontitis\textsuperscript{(28)}. These factors include: immunological host factors, ethnicity, microbiological factors, oral hygiene habits, age, gender, frequency of dental visits, demographic factors, smoking habits and psychological factors\textsuperscript{(29)}.

6. Histopathology and immunopathology

No major differences between aggressive and chronic periodontitis in terms of its histopathology and immunopathology are reported\textsuperscript{(1,2)}. Both appear as plasma-cell dominated lesions and mediated by Th2 cells. The localized form of aggressive periodontitis may represent a different entity with a genetic or epigenetic component. This could explain the association with the family history. On the other hand, aggressive generalized periodontitis could represent an advanced chronic periodontitis in young people with extreme
susceptibility, which would explain the common histopathological and immunopathological characteristics. These hypotheses need to be confirmed through rigorous studies\(^{(30)}\).

Localized aggressive periodontitis is frequently associated with deficiencies in the neutrophilic function and with high serum antibody response against periodontopathogens; while generalized aggressive periodontitis is also associated with deficiencies in neutrophil function, but with low serum antibody response against periodontopathogens\(^{(31)}\). Histopathological changes are reflected in the bone changes detected radiographically, even in mixed dentition\(^{(32)}\).

Furthermore, the colonization of the periodontal pocket by periodontal pathogenic bacteria could lead to an overlap of chronic periodontitis, which may complicate the histological and immunohistological condition\(^{(2)}\).

7. Microbiology

Some reports support the existence of subgingival microbiota resistant to antibiotics of choice\(^{(33-40)}\), which could explain eventual failures in the therapeutic modality. Localized aggressive periodontitis is mainly associated with the bacteria *Aggregatibacter actinomycetemcomitans*\(^{(41,42)}\), while generalized aggressive periodontitis is strongly associated with specific bacteria such as *Porphyromonas gingivalis*, *Tannerella forsythia*\(^{(43,44)}\) and *Aggregatibacter actinomycetemcomitans*, Gram-negative coccobacillus, capnophile, microaerophilic\(^{(45)}\). Microorganisms produce several virulence factors that could be involved in the destruction of periodontal tissues. The most important one seems to be leukotoxic activity\(^{(46,47)}\). The highly leukotoxic bacterial strains of Aa (strain JP2) can produce 10 to 20 times more toxins than other strains, giving them the potential to interfere with innate immune host defenses\(^{(48)}\). Some studies have shown that highly leukotoxic strains appear exclusively in individuals or families with a history of aggressive periodontitis\(^{(49-51)}\).

8. Clinical forms

8.1 Localized aggressive periodontitis

It begins at peripubertal age. It is mainly located in the first molars/incisors, with interproximal attachment loss in at least two permanent teeth, one of which is a first molar, and which
affects no more than two other teeth, apart from the first molars and incisors. It can also present atypical patterns, such as affecting other teeth instead of those mentioned.

8.2 Generalized aggressive periodontitis

It usually affects people under 30, but they may be older. There is an interproximal attachment loss which affects at least three permanent teeth additional to the first molars and incisors. Attached gingiva tissue loss is episodic\(^{(31)}\).

9. **DIAGNOSIS**

**Evaluation of the systemic condition**

The patient's medical history should be thoroughly evaluated. We must determine if there are risk factors such as smoking and psychosocial stress\(^{(5)}\). Since one of the characteristics of aggressive periodontitis is the absence of systemic diseases\(^{(32)}\), complementary tests can be run, if necessary, to rule out background pathologies. Record if the patient is taking any medication. Inquire about family history regarding periodontal condition\(^{(4)}\).

**Periodontal clinical examination**

The following should be evaluated: clinical attachment levels, periodontal pocket depth, bleeding on probing, furcations compromised, dental mobility, suppuration and oral hygiene.

**Radiological examination**

It is extremely important. The following are recommended:

- **Periapical radiographic series**: It should be done using the parallel technique and, preferably, a millimeter grid. There are the following options: seven radiographs for the upper arch and seven for the lower arch, two interproximal radiographs for molars and two for premolars.

- **Radiovisiography**: This is mainly indicated for cases of diagnostic doubt, regenerative therapy evaluation, periodontal status monitoring, among others. It is especially recommended for young patients with mixed dentition, where probing can be confusing. A distance greater than 2 mm between the cementoenamel junction and the alveolar crest in subjects with mixed dentition may suggest aggressive periodontitis.
Figure 1. Clinical photograph of 56-year-old female patient old with aggressive periodontitis.
A. Front view B. Side view.

10. MANAGEMENT ACCORDING TO COMPLEXITY LEVEL AND RESOLUTION ABILITY

Studies agree that treatment should be supplemented with antibiotics\(^{(53)}\). The objectives of the treatment are the same as for chronic periodontitis: reducing or eliminating the bacterial load and the contributory risk factors, in addition to regenerating the attachment apparatus as soon as possible.

11. TREATMENT PLAN SEQUENCE FOR PATIENTS WITH AGGRESSIVE PERIODONTITIS

11.1 Systemic Phase
- Medical referral, if indicated
- Medical interconsultation for the modulation of risk factors (tobacco, psychosocial stress)

11.2 Initial Phase
- Emergency treatment, if necessary
- Educating the patient about the disease process, contributing factors, perpetuating factors and triggers
- Teaching the patient about oral hygiene, evaluation and reinforcement of plaque control measures\(^{(4)}\)
- Study, diagnosis and treatment of occlusal disharmony and temporomandibular disorders\(^{(4)}\)
- Taking bacterial samples from selected pockets, cultures and antibiotic sensitivity testing can also be considered\(^{(4)}\)
- Prior dental treatments, if necessary
- **Supragingival and subgingival scaling and root planing.** Mechanical therapy is key in the treatment of aggressive periodontitis\(^{(4,53)}\). Even current consensus is that antimicrobial therapy should be preceded by mechanical debridement to break the structure of the biofilm\(^{(54)}\).
- Atraumatic extraction of non-viable teeth preserving the ridge.
- **Local and systemic antibiotic treatment.** The application of antibiotics via both pathways has advantages and limitations\(^{(54)}\).

**Systemic administration**

Prescribing systemic antibiotics for the treatment of aggressive periodontitis is properly supported\(^{(37, 38)}\). The combination of amoxicillin and metronidazole as an adjuvant is the best option, as described in recent systematic reviews\(^{(52,54-56)}\), especially for its effectiveness on *Aggregatibacter actinomycetemcomitans*\(^{(33)}\). Other recommended antibiotics for the treatment of aggressive periodontitis are metronidazole, spiramycin and clindamycin\(^{(34)}\). This method has the advantage of reaching all the areas of the oral cavity. However, optimal patient compliance is required to avoid an irregular administration that may not achieve the expected goals or may lead to bacterial resistance\(^{(54)}\).

**Local administration**

It allows for a high concentration of the drug in specific areas that would not be reached systemically; however, since not all the affected areas are treated, there may be reinfection\(^{(54)}\). Several studies show that an adjuvant therapy with local antimicrobials leads to pocket depth reduction and significant increases in the clinical attachment level compared
to the control groups. There is a better effect using tetracycline fibers, followed by doxycycline and minocycline. The application of chlorhexidine chips achieves minimal improvements in periodontal condition\(^{(54, 57, 58)}\).

11.3 Re-evaluation

- We should evaluate whether the treated sites show reduced probing depth, clinical attachment gain and resolution of inflammation. Oral hygiene and patient's motivation should also be evaluated\(^{(59)}\), in addition to tooth mobility, occlusal state and root sensitivity.

- Samples can be taken from selected pockets, medical interconsultation can be implemented, if indicated, and additional laboratory tests can be requested, if applicable.

11.4 Surgical Phase

Studies show that access or regenerative surgical treatments may have good results in patients with aggressive periodontitis, and that the effect may be similar to the ones obtained in chronic periodontitis; provided it is complemented with proper oral hygiene, risk factors are controlled and a rigorous maintenance program is followed\(^{(60)}\).

Resective, reparative or regenerative treatments can be carried out depending on the case. In aggressive periodontitis, antimicrobial surgical and comprehensive mechanical therapy is an appropriate treatment protocol for long-term stabilization of periodontal health\(^{(53)}\).

If there is improvement after surgical treatment, maintenance will be carried out; otherwise, we should return to the initial phase. It is important to monitor the surgically-treated areas.

11.5 Maintenance Phase

The success of the treatment will largely depend on patient compliance, both through adherence to doctor's instructions and by attending clinic appointments\(^{(56)}\). Medical appointments every three months have shown favorable results; however, based on the patient's periodontal status, the frequency may vary\(^{(55)}\).

- Monitor probing depth, clinical attachment level, gingival bleeding, tooth mobility\(^{(55)}\), patient's oral hygiene and risk factors\(^{(60)}\).
- Conduct annual radiographic examination of changes in the bone tissue of teeth at risk
- Perform scaling, root planing and polishing of residual pockets\(^{(55)}\)
- Topical fluoridation treatment and root desensitization (if indicated)
- Assess the need to resume the local or systemic antibiotic treatment
- Occlusal adjustment and balancing by selective wear of premature contacts in centric relation with or without pathological displacements (if necessary)
- Consider extracting teeth with terminal or unviable progressive disease using conservative and regenerative techniques of the residual alveolar bone.

**DIAGNOSIS**

- **Evaluation of systemic condition** (Discard diabetes, blood dyscrasia and other complications.) Consider the age of the patient.
- Inquire about family history of periodontal diseases. Find out the age at disease onset and if there has been tooth loss. Perform a stomatological clinical examination–periodontal chart. Determine how the loss progression of periodontal tissues occurred and if multiple teeth have been compromised.
- **Radiological analysis**: Vertical bone loss in proximal surfaces of posterior teeth. Several teeth may be affected. The pattern of bone loss is usually bilateral. Advanced cases may also present horizontal bone loss.

**TREATMENT**

**CONVENTIONAL INITIAL STAGE**
- Instruction and reinforcement for plaque control
- Bacterial monitoring of selected pockets
- Scaling and root planing
- Local and systemic antibiotic treatment
- Modulation of contributing local factors

**RE-EVALUATION**
- Evaluation of clinical attachment level
- Evaluation of periodontal probing depth
- Bacterial pocket monitoring
- Evaluation of risk factor modulation

**SURGICAL PHASE**
- Regenerative, restorative, resection or combined techniques, when indicated

**MAINTENANCE PHASE**
- Dental plaque control
- Evaluation of clinical attachment level, probing depth, gingival bleeding and tooth mobility
- Annual radiographs of parts at risk
- Host modulatory treatment
- Considering extracting unviable teeth

Inquire about family history of periodontal diseases. Find out the age at disease onset and if there has been tooth loss. Perform a stomatological clinical examination–periodontal chart. Determine how the loss progression of periodontal tissues occurred and if multiple teeth have been compromised.
Figure 2. Aggressive periodontitis diagnosis criteria flowchart (based on Albandar (2014)(32) and therapeutic routines for aggressive periodontitis

12. EXPECTED RESULTS AFTER AGGRESSIVE PERIODONTITIS TREATMENT:

- Significant reduction of clinical signs of gingival inflammation
- Reduction of probing depths
- Stabilization or gain of clinical attachment
- Radiographic evidence of resolution of bone lesions
- Reduction of clinically detectable plaque to a level compatible with an adequate level of periodontal health
- Progress toward occlusal stability(4,31)

The AAP(4) highlights the following considerations for these patients:

- Systemic evaluation of the patient to rule out diseases, hereditary traits and conditions that have periodontal repercussions
- The application of a mechanical treatment alone is usually not effective.
- The long-term treatment outcome will depend on the patient's care, compliance with maintenance appointments and plaque control skills. If primary teeth are affected, eruption of permanent teeth should be monitored to detect possible attachment loss.
- Due to the potential familial nature of aggressive diseases, evaluation and counseling of family members may be indicated(4).

It is common to fail in the treatment of aggressive periodontitis due to certain virulence factors of A. actinomycetemcomitans and P. gingivalis, which cannot be properly removed, due to their cell and tissue penetration ability(36).

CONCLUSIONS

Aggressive periodontitis are rare pathologies, mainly found in patients of African ethnicity or African descent. Early onset, family aggregation and rapid progression are usually their main characteristics. Consensus have been established on the forms of the disease: localized and
generalized. Immunological, genetic and microbiological factors are strongly associated, and seem to determine the two presentation forms. The concept of exclusivity of the periodontal pathogen \textit{Aggregatibacter actinomycetemcomitans} in aggressive periodontitis has been partially depreciated. Controversies have also arisen about the classification of aggressive periodontitis as a clinical entity independent from chronic periodontitis, because they share a common genetic basis, which would explain the phenotypic expression of the same disease. The diagnostic and therapeutic protocol does not differ much from that of other periodontal diseases; however special attention should be paid to family history, systemic condition, early diagnosis, specialized management by a periodontist, plaque control and frequent monitoring of periodontal pockets.

**REFERENCES**


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