## Case report

2	Dental considerations in a 4-year-old girl with Lennox-Gastaut Syndrome. Case report and
3	literature review.
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21	Abstract

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22 We present the developmental, oral, clinical, radiographic findings and oral treatment of a 4-

23 | year-old girl<u>presenting with</u> Lennox-Gastaut syndrome (LGS), which is a severe disabling

childhood epilepsy diseases that is treated with one or multiple anti-epileptic drugs (AEDs). The

25 child was wheel-chair bound, developmentally delayed, G-tube fed, and suffered from multiple 26 seizures and infantile spasms. The child's medical history included an under-developed pituitary gland, gastro esophageal reflux disease, vision and hearing impairment, history of chronic 27 28 aspiration pneumonia, and allergies. The oral findings included no carious lesions, heavy calculus accumulation, spontaneous bleeding from the gingiva, generalized gingival hyperplasia 29 (GH) and abnormal increased mobility in several primary teeth. The comprehensive radiographic 30 and clinical examination and the treatment under general anesthesia are described. The etiologies 31 of the calculus accumulation and GH are reviewed. 32

33 Key words: calculus, gingival overgrowth, anti-seizure medication

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## 35 <u>Introduction</u>

Lennox-Gastaut syndrome (LGS) is a severe and disabling childhood epilepsy that is 36 characterized by a triad of symptoms: 1) generalized treat seizures resistant to multiple therapies; 37 2) slowness of intellectual growth and cognitive impairment; 3) a specific electroencephalogram 38 (EEG) disturbance called a slow spike-and-wave pattern that is present when the child is 39 awake.<sup>1-5</sup> LGS patients may have multiple daily seizures that may cause sudden and 40 unpredictable stiffening followed by a drop to the ground; this being a key diagnostic feature.<sup>5-7</sup> 41 The pharmacologic treatment may include one or multiple antiepileptic drugs (AEDs),<sup>4</sup> some of 42 43 which have the potential to induce gingival hyperplasia (GH).

A review of the literature identified only one report of the oral findings in a LGS patient, of a
26-year-old female who had macroglossia, supragingival as well as subgingival calculus, red,
swollen and friable gingiva with generalized bleeding and localized suppuration, and gingival

47 recession.<sup>4</sup> The present manuscript includes an additional, comprehensive case report of a 448 year-old girl with LGS, and presents a review of the literature on LGS and related anti-seizure
49 medication that may have-induced gingival overgrowth.

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51 <u>Case presentation</u>

A 4.5-year-old Caucasian female with LGS was referred to a University Clinic for dental 52 treatment. The medical history indicated that she was born at 32 weeks of gestation, along with 53 her healthy twin. The patient had infantile seizures and spasms 15-16 times per day and was 54 diagnosed with LGS. Her medical history was significant for developmental delay, wheelchair-55 bound, had a gastrotomy tube (G-tube), under-developed pituitary gland, gastro esophageal 56 reflux disease, vision and hearing impairments, history of chronic aspiration pneumonia, 57 allergies to Depakote and Amoxicillin and leukodystrophy (degeneration of the white matter in 58 the brain<sup>8</sup>). Her medications included; Vigabatrin, Clobazam, Topiramate, Fycompa, Diazepam 59 and Rufinamide reducing the daily seizures to 3-6, and Albuterol/atropine via nebulizer. Recent 60 hospitalizations resulting from seizures, chronic pneumonia, and adrenocorticotropic hormone 61 therapy. The surgical history included adenoid and tonsils removal, Nissen fundoplication with 62 hernia repair, and G-tube placement. The chief complaint as expressed by her mother was risk of 63 aspirating exfoliating primary teeth: the previous night the patient had a seizure, after which she 64 was "choking and was missing a lower tooth that was swallowed or aspirated". 65

On examination, she had no apparent respiratory difficulties, was non-verbal, had a small
"hypoplastic" face, inability to cooperate, extensive drooling, short stature and slight
overweight.<sup>9</sup> A limited oral examination revealed sialorrhea, primary dentition with missing
mandibular primary central incisors, heavy calculus on the majority of teeth surfaces, abnormal

mobility (2-3 mm) in both mandibular primary lateral incisors (teeth #N and #Q), as well as
generalized moderate GH. Tongue size appeared normal. A chest radiograph did not reveal tooth
aspiration.

73 The patient MG was admitted to the hospital the day before the dental treatment under general anesthesia (GA), maintained with intravenous fluid to avoid the conflict between being 74 75 nil per os and her need for frequent G-tube feeding. The mother reported that the patient was 76 apparently having pain while grinding her teeth. Under GA, a radiographic and clinical examinations revealed no caries, no evidence of dental pulp pathology (Figure 1), all maxillary 77 78 primary incisors (Teeth D, E, F, G), and both mandibular lateral primary incisors (Teeth N and Q) had abnormal mobility ( about 3 mm), nearly all teeth were covered with heavy calculus 79 (Figure 2), generalized moderate GH, and a band of gingiva over the occlusal surface of the 80 mandibular right first primary molar (Tooth S, Figure 3A), and gingiva over the occlusal surface 81 82 of the maxillary right first primary molar (Tooth B, Figure 4a). The GH was non-hemorrhagic, soft, slightly fluctuant and pink (Figures 2, 3a&b, 4a&b). Calculus removal was accomplished 83 84 with an ultrasonic and hand instrumentation, followed by an application of a fluoride varnish. The gingival tissue over teeth B and S were removed with a surgical blade (Figures 3b & 4b). 85 Teeth # D, E, F, G, N and Q were extracted. The pot-operative recovery was uneventful. 86

## 87 Discussion and literature review

Dr. William Lennox, first described LGS in 1930s, Lennox and Davis later reported its
triad, which was further expanded by Gastaut.<sup>11-12</sup> The median onset age of LGS is about 4 years
(range: 0.6-28.9 years) with a peak onset of 5 years.<sup>13-14</sup> LGS is uncommon (3-10% of childhood
epilepsy) and has a mortality rate ranging from 3% to 7%.<sup>2, 3, 12</sup> The tonic seizures are
characterized by an EEG diffuse high voltage slow wave followed by generalized low voltage

fast activity, reflecting sustained fast neurological firing over a wide cortical area.<sup>5, 15</sup> 80% of
LGS patients will continue to have seizures into adulthood.<sup>2, 16</sup>

95 Based on our literature review, this is the second case in which the oral characteristics of 96 LGS are described, and the first one in a child. In our case the dental consideration included behavioral and management issues, gingival hyperplasia as a result of side effects caused by anti 97 98 seizure medication, poor oral hygiene (OH) and a risk of aspiration from loose teeth and difficulties in swallowing. Comparison of both cases is restricted by the patients different age 99 groups; the previous report was in a 26-year-old female.<sup>4</sup> Both cases received AEDs and had GH 100 101 and severe calculus accumulation, the previous case had periodontitis and macroglossia that 102 encumber proper OH while in the present case the tongue size was normal and there was increased abnormal tooth mobility. with no radiographic evidence of alveolar bone loss. Oral 103 pain was reported in the previous case associated with gingival swelling, gingival recession and 104 periodontitis while in the present case, pain was assumed to be related to biting on the gingival 105 tissue over the occlusal surfaces. 106 107 GH commonly starts with the eruption of the permanent dentition and may be influenced by genetic predisposition.<sup>17</sup> However, in the present case there was no history of GH in the 108 family,  $\frac{1}{10}$  indicating that the GH may have been caused by one or more AEDs most likely 109 vigabatrin. The aim of AEDs is to control or decrease seizures without producing unacceptable 110

adverse effects that impair quality of life; however, AEDs have been most frequently associated

with adverse drug reactions.<sup>17</sup> The pharmacologic treatment of LGS includes AEDs such as

113 vigabatrin, valproates, felmabate, and benzodiazepines which may potentiate each other<u>'s</u> side

114 effects, as in cases in which GH is potentiated by the combination of phenytoin and calcium

115 channel blockers, or cyclosporine and calcium channel blockers.<sup>18-20</sup>

Interestingly, multiple AEDs have an additive effect on GH, that might explain the additive
effect of multiple anticonvulsant therapy to GH.<sup>25</sup>

GH might include an abundance of dense connective tissue or acellular collagen that can be 118 an impediment to tooth eruption.<sup>36, 37</sup> Delayed eruption has also been associated with severe 119 bruxism in children with cerebral palsy.<sup>38, 39</sup> In the present case, the primary dentition was 120 normal.<sup>40</sup> However, the clinical crowns of the primary teeth appeared shorter than normal and 121 there was gingival tissue at the occlusal surfaces of teeth B and S, suggesting a combination of 122 GH and delayed eruption that could be related to the GH and bruxism (Figures ,3a, 3b). 123 Despite the positive correlation between plaque scores, gingival inflammation, and severity 124 of GH in children, the role of OH as an etiologic factor for GH has not yet been fully clarified 125 since most of the studies have been cross-sectional.<sup>19, 25</sup> However, the relevance of OH is 126

effective in controlling periodontal disease, and prevention of oral diseases is preferable for highrisk patients.<sup>4</sup> In the present case however, OH performance is complicated by the child's
inability to perform the most simple measures and to cooperate with her parents.

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emphasized in the previously reported LGS case in which non-surgical periodontal therapy was

A full mouth gingivectomy in the primary dentition was reported by Breen et al. (2009) in a case of a 28 month old with hereditary gingival fibromatosis in which only 4 mandibular teeth were partially erupted.<sup>17</sup> In the present case, we included the removal of the gingival tissue from the occlusal surfaces of the primary molars that most likely were the origin of oral pain (Figures 3b & 4b); in retrospective, a gingivectomy could have been adequate for the maxillary right primary cuspid and lateral incisor that had minimal clinical crowns (Figure 4a); the patient will continue to be under follow-up and will be scheduled for gingivectomy if required.

Children and adolescents who are unable to meet their nutritional needs orally and depend
on G-tube feeding at a significantly increased risk of poor oral health, specially tartar,
accumulation an subsequent gingivitis.<sup>10, 41, 46</sup> In the present case, the possibilities of recurrence
of calculus accumulation are high. Based on our search of the literature, it appears that this is the
youngest case reported with severe generalized calculus accumulation.

Aspiration of exfoliating primary teeth is apparently most uncommon or non-reported since 143 our review of the literature disclosed only oneuncommon. A case of aspiration of a maxillary 144 primary cuspid by a 9 year 11 month old child with cerebral palsy was reported, the authors 145 have -emphasized ing the fact that the possibility of aspiration of primary teeth is exacerbated in 146 debilitated patients.<sup>47</sup> Aalso, avulsion of primary teeth due to trauma and their aspiration is 147 possible.<sup>48</sup> This emphasizes the need to consider the need to investigate refer children who "lost" 148 a primary tooth that cannot be found tousing a chest radiograph, especially in children with 149 developmental disturbances, and a history of aspiration pneumonia which involves the entry of 150 infectious pharyngeal contents into the lower airway.<sup>41</sup> Relevant is the fact that low salivary flow 151 152 associated with gastric tube (GT) feeding may predispose the growth of salivary bacteria that, when mixed with food or liquid, provide a substantial inoculum to the lungs if aspirated.<sup>41</sup> 153 In conclusion, LGS in young child<u>ren</u> presents a significant challenge to the dental professional. 154 both Tthe nneurologist and the pediatric dentist should be aware of the potential complication and 155 work as team on behalf of the patient and the family of the LGS patient. 156

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276 Figure-2



- Figure-3b



283 Figure-4a



285 Figure-4b