3D stereophotogrammetry in children and adolescents with Scleroderma En Coup De Sabre/Parry-Romberg Syndrome: Description of a novel method for monitoring disease progression

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Abstract

Background: The diagnosis of Scleroderma En Coup De Sabre (ECDS)/Parry Romberg Syndrome (PRS) is mainly based on characteristic clinical findings. Methods to objectively monitor the course of the disease in a standardized way are lacking.

Objectives: This descriptive, retrospective, single centre cohort study aims to describe the contribution of 3D photographs in the assessment of the degree of facial asymmetry changes over time in growing children and adolescents with ECDS and PRS.

Methods: Six patients diagnosed with ECDS/PRS, with a follow-up period of at least 24 months and at least three 3D photographs were included. Mirroring these 3D photographs was automatically performed using surface-based matching to generate a colour-coded distance map, illustrating the inter-surface distance and thereby asymmetry between the original and mirrored 3D photographs. The percentage of absolute distances between the original and mirrored 3D photograph were calculated.

Results: In two patients, impressive decreases in the percentages of absolute distance levels over time were found, whereas the other patients did not show progression of asymmetry over time.

Conclusion: This study shows the potential of 3D stereophotogrammetry as an objective tool to measure disease activity over time in patients with ECDS/PRS.

1  |  INTRODUCTION

Morphea is an idiopathic sclerosing disorder of the skin and underlying tissues. A subtype of morphea, Scleroderma En Coup De Sabre (ECDS), affects the face and/or scalp with a linear induration of the skin and sometimes involvement of the underlying muscle and bone. Parry-Romberg Syndrome (PRS), also known as Progressive Hemifacial Atrophy, presents as a loss of tissue on one side of the face. It may involve dermis,
subcutaneous tissue, muscles and underlying bone. It is still controversial whether PRS is a variant of facial linear scleroderma or an independent disorder. Recent studies, however, have concluded that both diseases are on the same spectrum of disease, possibly sharing a similar pathogenesis. Therefore, some authors prefer to use the term linear scleroderma of the face to cover both ends of the spectrum.

When children and adolescents are affected by ECDS and/or PRS, the deformities potentially have influence on their facial attractiveness and psychological wellbeing. Currently, the diagnosis of ECDS/PRS is mainly based on characteristic clinical findings. The management of linear scleroderma of the face is challenging. Recently, minimum standards of care for children and adolescents with localized scleroderma (including scleroderma of the face) have been proposed by members of the Paediatric Rheumatology Europe Society Scleroderma Working Group. In their recommendations, the aim of treatment is to reach disease inactivity, defined as no erythema, no new lesions over the past 3 months, no worsening of skin thickness, no worsening of joint contractures, physician global assessment on visual analogue scale of 0 and no active extracutaneous involvement. The authors state that inactivity in deeper lesions can be harder to define by these parameters. Although erythema, induration and superficial extension of lesions can be assessed in ECDS and PRS, the deeper extension or progression of hemifacial atrophy is very difficult to monitor. Especially in growing children and adolescents who naturally show craniofacial growth, monitoring the disease is even more challenging. To the best of our knowledge, methods to objectively monitor the course of the disease in a standardized way are lacking.

To capture a facial soft tissue surface, three-dimensional (3D) stereophotogrammetry has gained great interest over the past decade in the field of Oral and Maxillofacial Surgery (OMFS). In patients who undergo maxillofacial orthognathic surgery, for example, 3D stereophotogrammetry is widely used to evaluate preoperative and postoperative three-dimensional changes of the face. The extra dimension of a 3D photograph makes it possible to calculate even the smallest surface differences. In addition, using surface-based registration techniques 3D photographs acquired at different stages of the disease and during treatment can be matched. This accurate quantification of the facial soft tissue surface is a validated method and can therefore be used to evaluate the progression of facial growth disorders over time. In case of disease activity of ECDS/PRS, deeper extension of skin lesions and/or progression of hemifacial atrophy is expected. We hypothesize that 3D photographs are able to accurately measure changes in facial asymmetry in ECDS/PRS and could therefore serve as a tool to monitor disease activity.

This study aims to describe the contribution of 3D photographs in the assessment of changes in facial asymmetry over time in growing children and adolescents with ECDS and PRS.

2 | MATERIAL AND METHODS

2.1 | Patient selection

In this descriptive, retrospective, single centre cohort study, patients diagnosed with ECDS/PRS treated at the Department of Paediatric Rheumatology and Dermatology at the Radboud University Medical Centre were selected. A few years ago, we decided to start making 3D photographs of all patients with ECDS/PRS visiting our combined outpatient clinic meaning that at the moment of the first 3D photograph, some patients already completed their systemic treatment while others were still on active drug or just commenced their systemic treatment. For this retrospective analysis, patients were included if they met the following inclusion criteria: (i) age at onset of ECDS/PRS <18 years, (ii) availability of at least three 3D photographs with an interval of 6 months during a follow-up period of at least 24 months. Patients who had (orthognathic) facial surgery during follow-up were not included.

This study was conducted in accordance with the World Medical Association Declaration of Helsinki on medical research ethics. The study was approved by the institutional medical ethical authority (file number W13_303 # 13.17.373) and informed consent was acquired for all patients who were enrolled in the study. All image data were anonymized and de-identified prior to analysis.
2.2 | Patient characteristics

The following items were retrospectively collected from medical electronic charts: age at onset of disease, age at first 3D photograph, age at inclusion in this study, sex, affected side and region of the face and previous and/or current medical treatment.

2.3 | Image acquisition

3D photographs were acquired using a stereo photogrammetrical camera set-up (3dMD face™ System, 3dMD LCC, Atlanta, GA, USA). All photographs were taken using a standardized protocol\(^{10}\): Patients were positioned in a natural head position with their eyes open, the forehead free of facial hair and their facial musculature in a relaxed state with loosely closed lips, in order to avoid overestimation of facial asymmetry.

2.4 | Analysis of facial asymmetry

To analyze facial asymmetry, all 3D photographs were processed as described by Verhoeven et al.\(^{13}\) In summary, four consecutive steps were needed to achieve a fine alignment of the 3D photographs (Figure 1). First, the region of interest was selected and confounding areas such as the neck were excluded. Second, a mirrored 3D facial surface (mirrored 3D photograph) was created using the midsagittal facial plane. For this step, an absolute perfect 3D photograph was necessary. Since our patients were in a growing phase, comparing 3D photographs from different time points was not possible. By creating a mirrored 3D photograph, patients were their own control group and a bias due to normal growth was excluded. After finishing this step an original and a created (mirrored) 3D photograph were available for detailed asymmetry analysis. Third, both original and mirrored 3D photographs were roughly aligned. After initial alignment, accurate surface-based alignment was automatically performed using surface-based matching (Iterative Closest Point algorithm). Last, a colour-coded distance map was generated to illustrate the inter-surface distance between the original and mirrored 3D photographs (Figure 2). Since there is great discussion of the mean facial asymmetry seen in the healthy paediatric and adolescent population, the obtained values were not corrected for this.\(^{14,15}\)

2.5 | Analysis of facial asymmetry over time

To analyze facial asymmetry over time for every individual, at least three colour-coded distance maps were created for the 3D photographs acquired at different timepoints using the abovementioned method. All patients included in the study had a follow-up period of at least 24 months, which made it possible to analyze potential progression of facial asymmetry.

2.6 | Statistical analysis

From the computed colour-coded distance maps the absolute mean value and standard deviations were calculated using MATLAB version R2019b (The Mathworks, Inc, Natick, Massachusetts, USA) for all patients and different timepoints. Descriptive analysis was performed using IBM SPSS Statistics, Version 25 (IBM Corp., Armonk, NY, USA).

3 | RESULTS

3.1 | Patient characteristics

In this descriptive pilot study six patients (50% females) were included (Table 1) with a median age at inclusion of 13 years [range 9–19 years] and a median duration of their disease of 5.5 years [range 5–8 years]. The median follow-up time from start of first 3D photograph to moment of analysis was 32 months [range 24–42 months].

Prior to the first 3D photograph, 5 out of 6 patients (83%) were treated with methotrexate (MTX). Of these
5 patients, 4 received this treatment for a period of at least 18 months [range 18–24 months] and one was treated with MTX for 12 months followed by mycophenolate mofetil (MMF) for a period of 15 months. Only one patient had the first 3D photograph taken before commencement of systemic therapy (patient 5). Apart from patient 2 and 3, all patients were still on MTX treatment when analysing the results.

### 3.2 Soft tissue asymmetry

Table 2 presents the absolute 3D distance measurements of the mirrored 3D photograph compared to the original 3D photograph.

TABLE 1 Patient characteristics

| Age at inclusion | Sex | Age of onset disease | Age at first 3D | Diagnose | Localization | Affected side | Treatment | Start treatment before first 3D photograph | Still on medication at moment of analysis |
|------------------|-----|----------------------|-----------------|----------|--------------|-------------|----------|------------------------------------------|------------------------------------------|
| Patient 1<sup>a</sup> | 11  | F                    | 5               | 8        | ECDS         | Forehead/chin | Left     | MTX                                     | Yes                                      | Yes                                      |
| Patient 2<sup>b</sup> | 19  | M                    | 11              | 16       | ECDS/PRS     | Forehead/chin | Left     | MTX                                     | Yes                                      | No                                       |
| Patient 3<sup>c</sup> | 16  | F                    | 11              | 13       | ECDS/PRS     | Mandible/Chin | Left     | MTX/MMF                                 | Yes                                      | No                                       |
| Patient 4<sup>d</sup> | 11  | F                    | 3               | 9        | ECDS/PRS     | Upper lip/eye | Left     | MTX                                     | Yes                                      | Yes                                      |
| Patient 5<sup>e</sup> | 14  | M                    | 9               | 12       | ECDS         | Forehead     | Left     | MTX                                     | No                                       | Yes                                      |
| Patient 6<sup>f</sup> | 9   | M                    | 4               | 7        | ECDS/PRS     | Mandible/chin | Left     | MTX                                     | Yes                                      | Yes                                      |

Abbreviations: ECDS, Scleroderma En Coup de Sabre; F, Female; M, Male; MMF, Mycophenolate Mofetil; MTX, Methotrexate; PRS, Parry-Romberg Syndrome.

<sup>a</sup>Patient 1 and 6 started MTX 2 years prior to the first 3D photograph, and are still on MTX for 24 and 42 months, respectively.

<sup>b</sup>Patient 2 received MTX for a period of 15 months, 15 months prior to the first 3D photograph. No medication during follow-up.

<sup>c</sup>Patient 3 received MTX for a period of 1 year and MMF for a period of 15 months prior to the first 3D photograph. No medication during follow-up.

<sup>d</sup>Patient 4 started MTX 2 years prior to the first 3D photograph, and is still on MTX for 24 months.

<sup>e</sup>Patient 5 received no medical treatment prior to the first 3D photograph, and is on MTX since first 3D photograph (24 months).

The percentage of absolute distances between the original and mirrored 3D photograph were calculated for cut-off values of 1 mm, 2 and 5 mm. These values describe which percentage of the face (3D photograph) lies within this range, with 100 indicating that all and 0 indicating that no surface points lay within the given range, respectively. In patient 2, for example, 36 months after baseline, only 44% of skin surface of the face lies within the cut-off value of 5 mm, compared to 92% at baseline. This indicates an absolute increase in asymmetry over time. The most impressive decreases in the percentages of absolute distance level less or equal in all three cut-off groups over time were noted in patients 2 and 3. Remarkably, only these two patients did not receive active drug therapy during the
period 3D photographs were taken. No clear difference in progression of asymmetry over time could be identified for the other patients.

4 | DISCUSSION

Linear scleroderma of the face is a rare condition mainly affecting children and adolescents. Since monitoring progression of hemifacial atrophy over time in an individual who naturally shows craniofacial growth is exceptionally challenging, the decision to quit or adjust medical treatment is mainly based on subjective parameters and expert opinions. To the best of our knowledge, a non-invasive method to objectively monitor the course ECDS and PRS over time in a standardized way is lacking. This study is the first to report that 3D photography can be used to objectively monitor disease progression over time in a growing individual.

3D stereophotogrammetry is known to be a safe and non-invasive imaging modality. It is a validated method widely used in OMFS which captures high quality 3D photographs of the soft tissue facial profile in less than 2 milliseconds without using radiation. Here we present a retrospective case series in which progression of facial asymmetry over time was found in two patients. Prior to their first 3D photograph, one patient (patient 2) was treated with MTX for 18 months, while another patient (patient 3) used MTX for 12 months and switched to mycophenolate mofetil due to gastro-intestinal complaints. Both patients refused further systemic treatment.
as they were satisfied with the clinical result and were reluctant to prolongate their treatment episode. The other four patients were still on MTX treatment at the moment of analysis (with a median treatment duration of 18 months) and showed no explicit progression of the facial asymmetry over time. These results might suggest a favourable effect of prolonged MTX treatment on the prevention of asymmetry. However, given the relatively small number of included patients, the descriptive nature of this study, lack of correction for confounding factors, solid conclusions on the efficacy of MTX on the prevention of facial asymmetry cannot be drawn.

As noted before, children and adolescents with facial asymmetry may suffer from negative psychological consequences. The degree of asymmetry causing these problems, however, is certainly not a predefined absolute value. In addition, every face has a slight degree of asymmetry. Cho et al. quantified a standardized normal craniofacial form and a baseline craniofacial asymmetry. They found a mean facial asymmetry in patients 0–18 years of age of 1.2 ± 0.6 mm. There was no statistical difference between age, sex, and race. These given values, however, are significantly higher in comparison with the values published by Kuijpers et al., who found an absolute mean facial asymmetry of 0.483 ± 0.148 mm. Since there is no consensus on the mean facial asymmetry of healthy children and adolescents, our results were not corrected for this. However, as the percentage of absolute distances between the original and mirrored 3D photograph in this study were calculated for cut-off values of 1, 2 and 5 mm, the percentages of 2 and 5 mm are absolutely indicative for a non-physiological facial asymmetry. A large prospective study is needed for this specific patient population to elucidate if this information can be used to adjust medical treatment and make clinical decision based on the progression measured using 3D stereophotogrammetry.

For clinicians, diagnosing severe facial asymmetry is not a challenge. The additional value of 3D photography is mainly in those patients showing mild progression of asymmetry over time. An overall progression of facial asymmetry of 1–2 mm between every consultation, for example, is very difficult to monitor. When using 3D photographs and the above-mentioned method to analyze facial asymmetry, the clinician is able to compare the current situation with the last check-up. With a system error of 0.2 mm and a reproducibility of 0.3–0.4 mm 3D stereophotogrammetry is able to capture small differences between these time-points. Indeed, the outcomes of the technique lead to changes in the treatment plan of our current daily clinical practice.

Di Giovanni et al. stated that cone beam computer tomography (CBCT) is also a reliable technique to assess bony as well as soft tissue changes (including the skin surface). There is a concern, however, that contraction of the facial muscles interferes with the absolute thickness of facial soft tissue measured with CBCT, potentially leading to a bias in the measured values. Second, one may question the clinical relevance of bony tissue evaluation of a young growing individual since reconstructive and aesthetic surgery is often advised after the face is fully developed. At last, even though it is less than a traditional CT-scan, CBCT still gives a radiation dose to the young patient.

The main limitations of the presented study are the limited number of included patients and the retrospective design. Since 3D stereophotogrammetry was a new imaging modality for patients with linear scleroderma of the face, 3D photographs of patients were collected at different stages of their disease and treatment phase. Further prospective research with a larger study population, 3D photographs prior to starting systemic treatment and a standardized protocol with a longer follow-up period is recommended to evaluate its use in monitoring progression of facial asymmetry. The main limitations of the 3D technique for daily clinical practice is the fact that disease activity on the scalp is difficult to monitor. Also, the currently used 3D photogrammetry technique is quite expensive, which makes it challenging to instantly implement it in every hospital. Comparing the outcomes of the currently used camera set-up with a less expensive handheld camera system would be interesting to investigate in further research in order to reduce costs.

In conclusion, this retrospective case series shows the potential of 3D stereophotogrammetry in patients with scleroderma of the face. This safe, validated and non-invasive imaging modality is able to detect changes in progression of asymmetry over time and may therefore be a useful tool to help clinicians to objectively monitor progression of linear scleroderma of the face over time in a growing individual.

**AUTHOR CONTRIBUTION**
Rutger ter Horst: Investigation, Writing – original draft. Thomas J. J. Maal: Software, Writing – review & editing. Martien J. J. de Koning: Supervision. Jorre S. Mertens: Supervision. Ellen J. H. Schatorjé: Writing – review & editing. Esther P. Hoppenreijis: Writing – review & editing. Marike M. B. Seyger: Supervision, Writing – review & editing.

**CONFLICTS OF INTEREST**
None to declare.

**DATA AVAILABILITY STATEMENT**
The data that support the findings of this study are available on request from the corresponding author. The data are not publicly available due to privacy or ethical restrictions.
ETHICS STATEMENT
This study was conducted in accordance with the World Medical Association Declaration of Helsinki on medical research ethics. The study was approved by the institutional medical ethical authority (file number W13_303 # 13.17.373) and informed consent was acquired for all patients who were enrolled in the study. All image data were anonymized and de-identified prior to analysis.

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REFERENCES
1. Zulian F. Scleroderma in children. Best Pract Res Clin Rheumatol. 2017;31(4):576–95.
2. Careta MF, Romiti R. Localized scleroderma: clinical spectrum and therapeutic update. An Bras Dermatol. 2015;90(1):62–73.
3. El-Kehdy J, Abbas O, Rubeiz N. A review of Parry-Romberg syndrome. J Am Acad Dermatol. 2012;67(4):769–84.
4. Tolkachjov SN, Patel NG, Tollefson MM. Progressive hemifacial atrophy: a review. Orphanet J Rare Dis. 2015;10:39.
5. Rhodes G. The evolutionary psychology of facial beauty. Annu Rev Psychol. 2006;57:199–226.
6. Kowner R. Psychological perspective on human developmental stability and fluctuating asymmetry: sources, applications and implications. Br J Psychol. 2001;92(part 3):447-69.
7. Constantin T, Foeldvari T, Pain CE, Pálnkás A, Hőger P, Moll M, Nemkova D, Weibel L, Laczkovski M, Clements P, Torok KS. Development of minimum standards of care for juvenile localized scleroderma. Eur J Pediatr. 2018;177(7):961-77.
8. Kau CH, Richmond S. Three-dimensional analysis of facial morphology surface changes in untreated children from 12 to 14 years of age. Am J Orthod Dentofacial Orthop. 2008;134(6):751–60.
9. Primozic J, Perinetti G, Contardo L, Ovensik M. Facial soft tissue changes during the pre-pubertal and pubertal growth phase: a mixed longitudinal laser-scanning study. Eur J Orthod. 2017;39(1):52–60.
10. Maal TJ, van Loon B, Plooj JM, Rangel F, Ettema AM, Borstlap WA, Bergé SJ. Registration of 3-dimensional facial photographs for clinical use. J Oral Maxillofac Surg. 2010;68(10):2391–401.
11. Laxer RM, Zulian F. Localized scleroderma. Curr Opin Rheumatol. 2006;18(6):606–13.
12. Dindaroglu F, Kutlu P, Duran GS, Görgülü S, Aslan E. Accuracy and reliability of 3D stereophotogrammetry: a comparison to direct anthropometry and 2D photogrammetry. Angle Orthod. 2016;86(3):487–94.
13. Verhoeven TJ, Coppen C, Barkhuysen R, Bronkhorst EM, Merx MAW, Bergé SJ, Mall TJJ. Three dimensional evaluation of facial asymmetry after mandibular reconstruction: validation of a new method using stereophotogrammetry. Int J Oral Maxillofac Surg. 2013;42(1):19–25.
14. Cho MJ, Hallac RR, Ramesh J, Seaward JR, Hermann NV, Darvann TA, Lipira A, Kane AA. Quantifying normal craniofacial form and baseline craniofacial asymmetry in the pediatric population. Plast Reconstr Surg. 2018;141(3):380e–7e.
15. Kuipers MA, Desmedt DJ, Nada RM, Bergé SJ, Fudalej PS, Maal TJ. Regional facial asymmetries in unilateral orofacial clefts. Eur J Orthod. 2015;37(6):636–42.
16. Lane C, Harrell W, Jr. Completing the 3-dimensional picture. Am J Orthod Dentofacial Orthop. 2008;133(4):612–20.
17. Boehnen CF, Patrick. Accuracy of 3D scanning technologies in a face scanning context. In: Proc. Fifth Int. Conf. 3-D Digital Imaging and Modeling; 2005.
18. Di Giovanni C, Puggina S, Meneghel A, Vittadello F, Martini G, Zulian F. Cone beam computed tomography for the assessment of linear scleroderma of the face. Pediatr Rheumatol Online J. 2018;16(1):1.

How to cite this article: ter Horst R, Maal TJJ, de Koning MJJ, Mertens JS, Schatorjé EJH, Hoppenreijis EP, et al. 3D stereophotogrammetry in children and adolescents with Scleroderma En Coup De Sabre/Parry-Romberg Syndrome: description of a novel method for monitoring disease progression. Skin Health Dis. 2022;2(3):e132. https://doi.org/10.1002/ski2.132