Production time, effectiveness and costs of additive and subtractive computer-aided manufacturing (CAM) of implant prostheses: A systematic review

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Abstract: OBJECTIVE To systematically review the dental literature for clinical studies reporting on production time, effectiveness and/or costs of additive and subtractive computer-aided manufacturing (CAM) of implant prostheses. MATERIALS AND METHODS A systematic electronic search for clinical studies from 1990 until June 2020 was performed using the online databases Medline, Embase and Cochrane. Time required for the computer-aided design (CAD) process, the CAM process, and the delivery of the CAD-CAM prostheses were extracted. In addition, articles reporting on the effectiveness and the costs of both manufacturing technologies were included. RESULTS Nine clinical studies were included reporting on subtractive CAM (s-CAM; 8 studies) and additive CAM (a-CAM; 1 study). Eight studies reported on the s-CAM of prosthetic and auxiliary components for single implant crowns. One study applied a-CAM for the fabrication of an implant bar prototype. Time was provided for the CAD process of implant models (range 4.9-11.8 min), abutments (range 19.7-32.7 min) and crowns (range 11.1-37.6 min). The time for s-CAM of single implant crown components (abutment/crown) ranged between 8.2 and 25 min. Post-processing (e.g. sintering) was a time-consuming process (up to 530 min). At delivery, monolithic/veneered CAD-CAM implant crowns resulted in additional adjustments chairside (51%/93%) or labside (11%/19%). CONCLUSIONS No scientific evidence exists on production time, effectiveness and costs of digital workflows comparing s-CAM and a-CAM. For both technologies, post-processing may substantially contribute to the production time. Considering effectiveness, monolithic CAD-CAM implant crowns may be preferred compared to veneered CAD-CAM crowns.

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Results: Nine clinical studies were included reporting on subtractive CAM (s-CAM; 8 studies) and additive CAM (a-CAM; 1 study). Eight studies reported on the s-CAM of prosthetic and auxiliary components for single implant crowns. One study applied a-CAM for the fabrication of an implant bar prototype. Time was provided for the CAD process of implant models (range 4.9–11.8 min), abutments (range 19.7–32.7 min) and crowns (range 11.1–37.6 min). The time for s-CAM of single implant crown components (abutment/crown) ranged between 8.2 and 25 min. Post-processing (e.g. sintering) was a time-consuming process (up to 530 min). At delivery, monolithic/veneered CAD-CAM implant crowns resulted in additional adjustments chairside (51%/93%) or labside (11%/19%).

Conclusions: No scientific evidence exists on production time, effectiveness and costs of digital workflows comparing s-CAM and a-CAM. For both technologies, post-processing may substantially contribute to the production time. Considering effectiveness, monolithic CAD-CAM implant crowns may be preferred compared to veneered CAD-CAM crowns.

Keywords
clinical research, clinical trials, material sciences, prosthodontics

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1 | INTRODUCTION

Computer-aided design (CAD) and computer-aided manufacturing (CAM) technologies used for the fabrication of implant-supported reconstructions enhanced time efficiency of the laboratory workflow compared to conventional manual techniques (Muhlemann et al., 2018; de Oliveira et al., 2020). Moreover, quality of implant prostheses fabricated by the aid of CAD-CAM technology meets the standards of prosthetic care (Mello et al., 2019; Muhlemann, Kraus, et al., 2018; Muhlemann et al., 2020).

Subtractive CAM (s-CAM) is considered the gold standard and referred to computer numeric controlled (CNC) milling of a desired shape out of a prefabricated material block. Limitations of this manufacturing process, however, apply. The waste of material, which includes also unused remnants of the blocks, may reach up to 90% (Strub et al., 2006). Fabrication accuracy applying s-CAM is influenced by the type of milling device (Bosch et al., 2014; Padros et al., 2020; Zeltner et al., 2017). Moreover, number, size and geometry of milling burs used for s-CAM seem to influence the surface resolution and represent a limitation that cannot be overcome (Atzeni & Salmi, 2012; Koch et al., 2016; Yara et al., 2005). Therefore, type and strategy of procedural milling determine the time for production, the quality of outcome and the costs for the milled prosthetic components.

Additive CAM (a-CAM) is referred to three-dimensional printing (3D printing) and is a process in which the desired object is produced by the deposition of layer upon layer (ISO/ASTM 52900-15, 2009). From an engineering point of view, a-CAM has the advantage to overcome the geometric restrictions that are present with s-CAM and more complex forms may be produced (Torabi et al., 2015). In addition, a-CAM may reduce waste of material (Zhang, Qiu, et al., 2019; Zhang et al., 2018).

Different a-CAM technologies exist and are applied in implant dentistry for processing polymers (Revilla-Leon & Ozcan, 2019), ceramics (Methani et al., 2020; Revilla-Leon et al., 2020) and metals (Revilla-Leon et al., 2019; Sun & Zhang, 2012). Applied a-CAM technology, processed materials, layer thickness and possible post-processing measures can affect the time of a-CAM procedures (Kaleli & Ural, 2020; Kessler et al., 2020; Presotto et al., 2019; Sames et al., 2016; Silva et al., 2011).

It is often postulated that a-CAM is a time-efficient and cost-effective manufacturing technology (Revilla-Leon et al., 2020; Williams et al., 2020). There is no general consent, however, to support or reject the potential advantages of a-CAM technologies compared to the s-CAM technologies in terms of fabrication efficiency and effectiveness for fixed and removable implant prostheses. Therefore, the aim of the present systematic review was to systematically assess the dental literature reporting on production time, effectiveness and/or costs of CAD-CAM implant prostheses involving s-CAM and a-CAM.

2 | MATERIAL AND METHODS

2.1 | Protocol development and eligibility criteria

A protocol was developed and registered in PROSPERO (CRD42020195942). Conducting and reporting was performed according to the Preferred Reporting Items for Systematic Review and Meta-analyses (PRISMA) statement (Liberati et al., 2009; Moher et al., 2009).

2.2 | Focused question

What are the production time, the effectiveness and costs for s-CAM and a-CAM technologies applied in the digital workflow for the fabrication of fixed and removable implant prostheses?

2.3 | PICO

The following PICO terms were used:

P Population: patients receiving fixed or removable implant prostheses
I Intervention: fabrication of fixed or removable implant prostheses applying a-CAM technologies
C Comparison: fabrication of fixed or removable implant prostheses applying s-CAM
O Outcome: production time (primary outcome), effectiveness (number of prostheses in need of chairside/laboratory adjustments at delivery) and costs (secondary outcomes)

2.4 | Search strategy

Three online databases Medline (PubMed), Embase and Cochrane Central Register of Controlled Trials (CENTRAL) were screened for eligible studies. The search was limited to clinical studies published from 1 January 1990 to June 2020 in the dental literature and in English language. An additional hand search was performed by screening the reference list of all included full-text articles.

2.4.1 | Search protocol

The following search terms were used:

For identifying the “population”:

Implants
[MeSH terms]: Dental implants OR Dental Implants, Single-Tooth OR Dental Implantation OR Dental Implantation, Endosseous OR
[Text Words]: “implant***” OR
[Entree terms]: tooth implant OR single tooth implant OR tooth implantation
Reconstructions/dentures
[MeSH terms]: Dental Prosthesis OR Dental Prosthesis, Implant-Supported OR denture, fixed partial OR dentures, fixed partial OR Crowns OR Dentures OR Overdenture OR Overdentures OR
Dental restoration, Permanent OR Tooth, Artificial OR Dental abutments
OR
[Text Words]: "prosth*" OR "replacement*" OR "reconstruction*" OR "restoration*" OR "suprastructure*" OR "restoration" OR "crown*" OR "fixed dental prosthes*" OR "fixed partial denture*" OR "bridge*" OR "full-arch*" OR "framework*" OR "bar*" OR "denture*" OR "abutment*" OR "attachment"
OR
[Emtree terms]: tooth prosthesis OR implant-supported denture OR tooth crown OR fixed partial denture OR fixed dental prosthesis OR denture OR overlay denture OR dental restoration OR dental abutment
For identifying the "intervention/comparison":

[MeSH terms]: Dental Technology OR Computer-Aided Design OR Computer-Aided Manufacturing OR Manufacturing, Computer Aided OR Design, Computer Aided OR Printing, Three Dimensional OR Printings, Three-Dimensional OR Three-Dimensional Printings OR 3-Dimensional Printing OR 3 Dimensional Printing OR 3-Dimensional Printings OR Printing, 3-Dimensional OR Printings, 3-Dimensional OR 3-D Printing OR 3 D Printing OR 3-D Printings OR Printing, 3-D OR Printings, 3-D OR Three Dimensional Printing OR Three Dimensional Printing OR 3D Printing OR 3D Printings OR Printing, 3D OR Printing, 3D OR Stereolithography OR Hot Melt Extrusion Technology OR
[Text Words]: "digital" OR "virtual" OR "intraoral impression" OR "IOS" OR "scan" OR "techn*" OR "CAD" OR "CAM" OR "computer" OR "manufactur*" OR "design" OR "fabricat*" OR "producing*" OR "additive" OR "print*" OR "3D" OR "stereolithography" OR "SLA" OR "digital light processing" OR "direct light processing" OR "DLP" OR "material jetting" OR "MJ" OR "direct inkjet printing" OR "inkjet printing" OR aerosol jet printing OR "material extrusion" OR "ME" OR "fused deposition of ceramic" OR "FDM" OR "multiphase jet solidification" OR "MJS" OR "extrusion free forming" OR "solid free forming" OR "EFF" OR "sintering" OR "direct ink writing" OR "DIW" OR "direct-write assembly" OR "DWA" OR "microrobotic deposition" OR "μRD" OR "three-dimension fibre deposition" OR "micropen" OR "power bed fusion" OR "PBF" OR "selective laser sintering" OR "SLS" OR "powder" OR "slurry coating" OR "aerosol-assisted spray deposition" OR "selective laser melting" OR "SLM" OR "slurry spraying" OR "ring blade" OR "direct energy deposition" OR "direct laser cladding" OR "hybrid fused deposition modeling" OR "sheet lamination modelling" OR "laminated object modelling" OR "LOM" OR "traditional sheet lamination" OR "sheet lamination" OR computer-aided manufacturing of laminated engineering materials" OR "binder jetting" OR "3D printers" OR binder jetting of dry powder agglomerates" OR Slurry-based 3D printing” OR "subtractive" OR "mill*" OR "cut*" OR "CNC" OR "trajectory" OR "bur"
OR
[Emtree terms]: clinical technology OR computer aided design OR computer aided design/computer aided manufacturing OR dental CAD/CAM system OR stereolithography OR three-dimensional printing OR fused deposition modeling OR selective laser sintering OR hot melt extrusion OR robocasting OR powder OR selective laser melting OR fused deposition modelling OR digital light processing OR electrophoretic deposition OR milling
For identifying the "outcome":

[MeSH terms]: Clinical Effectiveness OR Effectiveness, Clinical OR Treatment Effectiveness OR Effectiveness, Treatment OR Treatment Efficacy OR Clinical Efficacy OR Efficacy, Clinical OR Time Management OR Cost Effectiveness OR Effectiveness, Cost OR Cost Benefit OR Costs and Benefits OR Benefits and Costs OR Cost-Effectiveness Analysis OR Analysis, Cost-Effectiveness OR Cost Effectiveness Analysis
OR
[Text Words]: "efficiency*" OR "effect*" OR "time*" OR "effort*" OR "cost*" OR "money*" OR "finance*" OR "economic*" OR "deliver*" OR "remake*" OR "chairside*" OR "adjust*" OR "intervention*"
OR
[Emtree terms]: productivity OR clinical effectiveness OR therapy OR time OR time management OR exercise OR cost OR cost effectiveness analysis OR money OR finance OR economic aspect OR dental CAD/CAM system

2.4.2 | Final search strategy
(Implants AND Reconstructions/Dentures) AND (Intervention/Comparison OR outcome).

2.4.3 | Inclusion criteria
Clinical studies with a minimal number of five patients and studies conducted in the dental laboratory with at least five clinical cases were included.

2.5 | Exclusion criteria
In vitro and preclinical studies, interviews, charts, case reports with less than five patients and reports based on questionnaires were excluded from this systematic review. Studies on provisional or interim prostheses were not included.

2.6 | Selection of publications
Two reviewers (SM and JH) performed the screening independently. Titles and abstracts were evaluated for suitability. In case of disagreement, the decision was made by discussion between all authors.
Full-text articles of selected abstracts were acquired, and the final selection based on inclusion/exclusion criteria was made. For the final inclusion, the Material and Methods, Results and Discussion of the full-text articles were assessed and double-checked by two reviewers (SM and JH). Again, in case of disagreement during the evaluation, consensus was attained by discussions between all authors. As a measure of agreement, Cohen's kappa coefficient was calculated for abstract and full-text screening.

### 2.7 Data extraction and method of analysis

The following parameters were extracted and recorded from the selected full texts (Table 1): author(s), year of publication, study design, number of patients/cases, mean age, implant system, prosthesis design (fixed, removable), number of prostheses, type and system of digitalization (conventional/IOS), CAD system, CAM system, type of CAM (subtractive/additive), location of CAM and finalization of prosthesis (point-of-care, dental laboratory). Authors were contacted for clarification in case the included studies were lacking information to properly extract the data.

### 2.8 Evaluation of quality

The methodological quality of the included studies was assessed independently by two reviewers (SM and JH) using Cochrane Collaboration's tool for assessing risk of bias (Higgins & Green, 2011). For non-randomized studies, modified risk assessment tool was used. In case of disagreement, the decision was made by discussion between all authors.

#### TABLE 1 Description of included studies

| Author/Year | Study design | Subjects | Mean age | Implant system | Design prosthesis | Number of prosthesis |
|-------------|--------------|----------|----------|----------------|------------------|----------------------|
| Di Fiore et al. (2018) | prospective cohort study with a crossover design | 10 | na | na | fixed | 10 |
| Joda and Bragger (2014) | prospective cohort study | 6 | na | Straumann Tissue Level | fixed | 3 |
| Joda and Bragger (2014) | prospective cohort study with a crossover design | 20 | 55.4y | Straumann Tissue Level | fixed | 20 |
| Joda and Bragger (2016) | Randomized controlled trial | 20 | 55.4y | Straumann Tissue Level | fixed | 10 |
| Mangano, Margiani, et al. (2019) | retrospective cohort study | 25 | 51.1 | Exacone, Leone Implants | fixed | 40 |
| Mangano and Veronesi (2018) | randomized controlled clinical trial | 50 | 52.6 | Exacone, Leone Implants | fixed | 25 |
| Mangano, Mangano, et al. (2019) | prospective cohort study | 15 | 68.8 | BTSafe, BTK | removable | 15 |
| Pan et al. (2019) | prospective cohort study with a crossover design | 40 | 45.1y | Straumann Tissue Level and Bone Level | fixed | 40 |
| Zhang, Qiu, et al. (2019) | Randomized controlled trial | 33 | 46.8y | CAMLOG SCREW-LINE, Camlog Biotechnologies AG | fixed | 17 |
3  |  RESULTS

3.1  |  Search

The details of the search strategy are illustrated in Figure 1. The electronic search identified a total of 7,482 titles. After the evaluation of titles, 7,219 studies were discarded and 20 duplicates were removed. Following the screening of 243 abstracts, 34 studies were selected for detailed reading of the full text (Cohen’s kappa coefficient = 0.78). Finally, 8 studies met the inclusion criteria (Cohen’s kappa coefficient = 0.72). One study was added through the manual search, resulting in an overall number of 9 included articles.

The reasons for exclusion of studies are depicted in Table S1: description of a CAM process without information on time and/or effectiveness (n = 14), no detailed data on fabrication method (1), no CAM fabrication (1), no implant reconstruction (n = 1), outcomes limited to clinical follow-up (6), interview (n = 1), interim prosthesis (n = 1), and full text not in English language (n = 1).

3.2  |  Description of studies

All characteristics of the included studies are described in Table 1. Eight included studies applied s-CAM technologies for the fabrication of single implant crowns in the posterior area. One included study used a combination of a-CAM and s-CAM technologies for the fabrication of an implant bar serving as retention for a maxillary overdenture. No studies were identified reporting on the CAM fabrication of multi-unit fixed implant prostheses considering the primary and secondary outcomes.

None of the three randomized controlled clinical studies was primarily assessing production time, effectiveness and costs comparing
s-CAM and a-CAM. Consequently, a total of 9 prospective clinical studies reporting on 11 patient cohorts were identified. Five studies reported on the production time using s-CAM technologies for the fabrication of models (2), customized abutments (3), monolithic crowns (4) and crown cores (2). Time for the CAD process was reported in all five studies, whereas the time for the manufacturing
## TABLE 2  Primary and secondary outcomes of included studies (na, not applicable)

| Study                          | Manufacturing time                                                                 | Effectiveness                                      | Costs                               |
|-------------------------------|-----------------------------------------------------------------------------------|----------------------------------------------------|-------------------------------------|
| Di Fiore et al. (2018)        | n.a.                                                                              | Time for chairside adjustments                      | n.a.                               |
| Joda and Bragger (2014)       | n.a.                                                                              | Number of crowns in need of chairside               | Total laboratory costs              |
| Joda and Bragger (2015)       | CAD model<br>CAD customized abutment<br>CAD crown coping<br>n.a.<br>Ceramic veneering<br>glazing and polishing | Time for chairside adjustments                      | n.a.                               |
| Joda and Bragger (2016)       | CAD model<br>CAD monolithic crown<br>CAD customized abutment<br>n.a.<br>Assembly of components<br>polishing<br>Veneering<br>glazing and polishing | Time for chairside adjustments                      | Total laboratory costs              |
| Mangano, Margiáni, et al. (2019) | n.a.                                                                              | Number of crowns in need of chairside and laboratory adjustments | n.a.                               |
| Mangano and Veronesi (2018)   | CAD customized abutment<br>CAD monolithic crown<br>Subtractive<br>Sintering assembly of components<br>staining | n.a.                                               | Customized abutment<br>Monolithic crown<br>staining |
| Mangano, Mangano, et al. (2019) | n.a.                                                                              | Number of 3D-printed prototype bars with passive fit | n.a.                               |
| Pan et al. (2019)             | CAD monolithic crown<br>n.a.<br>n.a.                                               | Time for chairside adjustments<br>Number of crowns in need of chairside and laboratory adjustments | n.a.                               |
| Zhang, Qiu, et al. (2019)     | CAD monolithic crown<br>CAD crown coping<br>subtractive<br>Fitting and staining<br>Sintering veneering | Time for chairside adjustments<br>Number of crowns in need of chairside and laboratory adjustments | n.a.                               |
process was evaluated in two and the post-processing/finalization of the prostheses in 4 out of these five studies (Table 2). One study reported on the total working time applying a s-CAM process without further specification and details. Therefore, that study was excluded from the analysis (Joda & Bragger, 2014). Effectiveness and fabrication costs (secondary outcomes) were reported in 8 and 3 of the included studies, respectively.

3.3 | Risk of bias in included studies

Table 3 describes the risk of bias assessment of the nine included studies. For the double-blind self-controlled clinical study (Pan et al., 2019), the risk of bias was rated low in all categories, whereas for the three randomized controlled clinical trials, a high risk of performance bias and an unclear risk of detection bias were estimated. Considering the primary and secondary outcomes, attrition bias was rated low in all included studies.

3.4 | Primary outcome: Production time

3.4.1 | Digital workflow applying centralized CAM

In three studies, centralized s-CAM was involved, whereas the CAD process and the finalization of the prostheses were based in the dental laboratory. Two studies reported on a calibrated workflow (Straumann CARES) using a laboratory CAD software and centralized s-CAM provided by the implant manufacturer (Joda & Bragger, 2015, 2016), while one study used a laboratory CAD software (3Shape Designer) and an unspecified third-party provider for centralized s-CAM (Zhang et al., 2019).

The time for the virtual design (CAD) of (a) an implant model ranged between 4.9 min (Joda & Bragger, 2015) and 11.8 min (Joda & Bragger, 2016), (b) an abutment from 19.7 min (Joda & Bragger, 2016) up to 32.7 min (Joda & Bragger, 2015), (c) a monolithic crown accounted for 22.3 min (monolithic crown) (Joda & Bragger, 2016) and (d) a crown coping between 11.1 min (Zhang, Tian, et al., 2019) and 37.6 min (Joda & Bragger, 2015) (Table 4).

Production time for the centralized s-CAM process was reported in one study (Zhang, Tian, et al., 2019). The milling of the crown coping (zirconia) accounted for 8.2 min and the sintering lasted for 530 min. No information on delivery times of these centrally manufactured products to the dental laboratory was available.

The time needed for the finalization of the CAM products in the dental laboratory depended on the type of prosthesis. The veneering was most time-consuming and accounted for 71.4 min (Joda & Bragger, 2015), 74.7 min (Joda & Bragger, 2016) and 89.9 min (Zhang, Tian, et al., 2019). Glazing/polishing ranged between 11.5 min (Joda & Bragger, 2015) and 28.8 min (Joda & Bragger, 2016). For the monolithic design, the assembly of the prosthetic components took 5.0 min followed by the polishing procedure accounting to 15.4 min (Joda & Bragger, 2016).
| Preparation task | Time (min) | s-CAM | Time (min) | Post-processing | Time(min) |
|-------------------|------------|-------|------------|-----------------|-----------|
| **Digital workflow applying centralized s-CAM** | | | | | |
| Joda and Bragger (2015) | CAD model | 4.9 | n.a. | Ceramic Veneering | 71.4 |
| | CAD customized abutment | 32.7 | | glazing and polishing | 11.5 |
| | CAD crown coping | 37.6 | | | |
| Joda and Bragger (2016) | CAD model | 11.8 | n.a. | Assembly of components | 5.0 |
| | CAD monolithic crown | 22.3 | | Polishing | 15.4 |
| | CAD customized abutment | 19.7 | | Veneering | 74.7 |
| | | | | Glazing and polishing | 28.8 |
| Zhang, Qiu, et al. (2019) | CAD crown coping | 11.1 | Milling zirconia crown coping | Sintering | 530.0 |
| | | | | Veneering | 89.9 |
| **Digital workflow applying laboratory s-CAM** | | | | | |
| Mangano and Veronesi (2018) | CAD customized abutment | 30.0 | Milling zirconia abutment/crown | Sintering | 480.0 |
| | CAD monolithic crown | 10.0 | assembly of Components | 10.0 |
| Pan et al. (2019) | CAD monolithic crown | 12.6 | n.a. | n.a. | |
| **Digital workflow applying chairside s-CAM** | | | | | |
| Zhang, Qiu, et al. (2019) | CAD monolithic crown | 28.9 | Milling lithium disilicate crown | Fitting and staining | 40.2 |
3.4.2 | Digital workflow applying laboratory CAM

In two studies, the entire fabrication process was located in the dental laboratory. The laboratory CAD software was applied for the virtual design of a monolithic crown (3Shape Designer) (Pan et al., 2019), and an implant crown consisting of a customized abutment and a monolithic crown (Exocad DentalCAD) (Mangano & Veronesi, 2018).

The time for the CAD of a monolithic zirconia crown ranged between 10.0 min (Mangano & Veronesi, 2018) and 12.6 min (Pan et al., 2019), while the customization of the abutment took 30.0 min (Mangano & Veronesi, 2018) (Table 4).

The time for the s-CAM of the zirconia abutment/crown by means of a 5-axis milling machine (Roland DWX-50) was 25 min followed by 480 min of sintering (Mangano & Veronesi, 2018). In the second study, the manufacturer of the laboratory-based s-CAM device was provided (Zenotec Select Hybrid), but no time recordings were performed (Pan et al., 2019).

The final assembly of the hybrid abutment (cementation of customized zirconia abutment to titanium base) and the characterization of the monolithic crown was 10 min each (Mangano & Veronesi, 2018).

3.4.3 | Digital workflow applying chairside CAM

One study reported on a point-of-care CAD-CAM system (Cerec) in a chairside workflow (Zhang, Tian, et al., 2019). The CAD of a monolithic crown took 28.9 min and the s-CAM of lithium disilicate by means of a point-of-care milling device (Cerec MC XL, Sirona Dentsply) was 21.4 min. Post-processing included fitting and staining and amounted to a total of 40.2 min (Table 4).

3.5 | Secondary outcomes

3.5.1 | Effectiveness

Five studies reported on the effectiveness for monolithic implant crowns fabricated by means of s-CAM (Di Fiore et al., 2018; Joda & Bragger, 2016; Mangano et al., 2019; Pan et al., 2019; Zhang, Tian, et al., 2019)). At delivery, the time for chairside adjustments ranged from no intervention needed (Joda & Bragger, 2016) to 1.9 min (Di Fiore et al., 2018), 8.4 min (Zhang, Tian, et al., 2019), up to a range between 11.4 and 12.3 min (Pan et al., 2019). Three studies reported time for adjustments at delivery of veneered implant crowns consisting of CAM products. The chairside time ranged between 2.6 min (Joda & Bragger, 2015), 3.3 min (Joda & Bragger, 2016) and 12.3 min (Zhang, Tian, et al., 2019).

Five studies provided frequency distributions on the number of monolithic implant crowns in need of chairside or laboratory adjustments (Joda & Bragger, 2014, 2016; Mangano, Margiani, et al., 2019; Pan et al., 2019; Zhang, Tian, et al., 2019). Out of a total of 136 (96 model-free and 40 model-based) laboratory-made monolithic implant crowns, 50.7% (range 0% to 80%) and 11.0% (range 0%–13.8%) needed chairside adjustments and laboratory interventions (redo or adjustment), respectively. Of model-free crowns, 43.8%/10.4% needed chairside/laboratory adjustments, whereas of model-based crowns, 70%/12.5% needed chairside/laboratory adjustments. In a chairside workflow, 15 out of 17 (88.2%) monolithic lithium disilicate implant crowns were adapted chairside to be delivered, whereas one crown needed to be redone in a subsequent laboratory workflow (Zhang, Tian, et al., 2019). For veneered CAD-CAM implant crowns, 24/26 (92.3%) needed chairside interventions and additional laboratory veneering was needed in 5/26 (19.2%) (Joda & Bragger, 2016; Zhang, Tian, et al., 2019).

One clinical study evaluated the effectiveness of the fabrication of a milled polyether-ether-ketone (PEEK) implant bar fixed on 4 implants in the maxilla (Mangano et al., 2019). The clinical protocol involved the try-in of a 3D-printed resin bar and demonstrated that 3 out of 15 (20%) resin bars did not have a sufficient fit or adaptation. In these 3 cases, data acquisition and processing were repeated. Finally, all milled PEEK bars could be successfully delivered without remake.

3.5.2 | Costs

Laboratory costs for implant crowns were documented in three studies. Based on a laboratory tariff system, the total costs ranged between CHF 505.85 (Joda & Bragger, 2016) and CHF 785 (Joda & Bragger, 2014) for monolithic implant crowns. One study provided detailed costs for the CAM products: EUR 50 for a hybrid abutment, EUR 80 for a monolithic zirconia crown and additional EUR 30 for the characterization of the crown (Mangano & Veronesi, 2018).

4 | DISCUSSION

The present systematic review revealed (a) production time, effectiveness and costs to be thoroughly documented for fabrication workflows involving s-CAM, (b) production time for the implant crowns to range substantially depending on the prosthesis design and the applied digital workflow, (c) the laboratory fabrication of monolithic implant crowns to result in the least number of chairside and labside interventions and (d) a lack of clinical scientific evidence on production time for fabrication workflows applying a-CAM.

A recent systematic review demonstrated the implementation of digital technologies to substantially increase time efficiency in the laboratory fabrication of implant prostheses compared to manual techniques (Mühlemann, Kraus, et al., 2018). The present systematic search provides data on time required for all steps of the digital workflow involving s-CAM technology including the CAD process, the CAM production and the laboratory finalization of the CAD-CAM products. In addition, time for s-CAM is documented in chairside (point-of-care), laboratory and centralized workflows.
No clinical study, however, evaluated production time for digital workflows involving a-CAM. Therefore, it remains unclear whether fabrication workflows with a-CAM are more efficient compared to workflows with s-CAM.

Time for CAD varied greatly among the included studies. In general, the number of prosthetic/auxiliary components and the need for customization influenced production time. The calculated time differences for the CAD process among the included studies were further affected by (a) the heterogeneity of the study design, (b) the applied software systems and (c) the software version (Haddadi et al., 2018; Shim et al., 2015). The digital systems are constantly improving and, for example, resulted in a 70% more efficient CAD process for a monolithic implant crown between 2015 (37.6 min) (Joda & Bragger, 2015) and 2019 (12.0 min) (Pan et al., 2019).

The restoration materials processed by s-CAM were zirconia and lithium disilicate (Mangano & Veronesi, 2018; Zhang, Tian, et al., 2019). Production time of single implant components by means of s-CAM ranged between 8.2 and 25 min. Time differences may be attributed to (a) the different s-CAM systems applied (e.g. 4-axis versus 5-axis milling devices) and (b) the different restoration materials processed.

The present systematic review revealed that post-processing is a time-consuming process. Sintering of zirconia may take more than 8 hr (Mangano & Veronesi, 2018) compared to lithium disilicate accounting for 40.2 min (Zhang, Tian, et al., 2019). Consequently, zirconia is not the material of choice for chairside restorations. Recent in vitro studies, however, demonstrated that speed sintering of zirconia may allow to overcome this limitation while still maintaining its favourable mechanical properties (Jerman et al., 2020; Michailova et al., 2020).

The costs of CAD-CAM technology are rather high when data are compared to manual fabrication in the dental laboratory (Mühlemann, Kraus, et al., 2018). For example, one included study reported on the costs of prosthetic components (between EUR 50 and 80) processed by means of s-CAM (Mangano & Veronesi, 2018). It remains unknown though whether these costs covered all expenses including the salary of the operator, the restoration material, any consumables (e.g. milling burs) and the costs for the amortization of the CAM device. Still, the possibility of using services offered by external parties may allow accessing cost-effective CAD-CAM products.

None of the included studies reported on the production time when using a-CAM. Until recently, a-CAM technologies were too expensive to be applied in the fabrication workflow of implant prostheses, thereby explaining the limited number of clinical studies. Moreover, the development of materials (a) to be processed by means of a-CAM and (b) to be successfully applied in implant prosthetics is still in its early phase (Revilla-Leon et al., 2019; Revilla-Leon, Meyer, et al., 2020; Revilla-Leon & Ozcan, 2019).

The production time by means of a-CAM depends on the respective technology and the printer settings applied. Generally, an increase in layer thickness results in a faster production of the object (prosthetic or auxiliary component). Such an increase in layer thickness (i.e. a faster production) negatively affects the surface resolution (Kaleli & Ural, 2020; Kaleli et al., 2019) and the mechanical properties of the restoration (Shim et al., 2020). One has to bear in mind that the capacity of the a-CAM technology (Kessler et al., 2020) and the properties of the processed a-CAM material (Sames et al., 2016) may limit the layer thickness, thereby controlling the production time. Finally, additional programming (nesting) may be indicated for an ideal orientation of the virtual object relative to the printing direction to improve the marginal and internal adaptation of the prosthesis (Shim et al., 2020).

A-CAM involves post-processing procedures. The removal of supporting structures depends on the a-CAM technology applied and consequently influences the production time. In case of the material jetting technology, the co-deposited uncured material serves as a supporting structure and its removal is achieved by a fast cleaning process. In contrast, complex supporting structures are crucial to prevent a collapse of the object during the manufacturing process when stereolithography (SLA) or digital light processing (DLP) technologies are applied (Presotto et al., 2019; Silva et al., 2011). Moreover, the object (prosthetic/auxiliary component) can be chemically and/or mechanically treated, and curing cycles may be added. The need, type and time for post-processing procedures depend on the a-CAM technology and the manufacturer’s recommendations.

The thorough search performed in the present systematic review resulted in only one clinical study applying a digital workflow involving a-CAM. A-CAM technology was used for the fabrication of an auxiliary component as part of the fabrication workflow for an implant-retained bar (Mangano, Mangano, et al., 2019). A prototype bar was 3D-printed using a polymer with the intent to increase the effectiveness of a workflow using s-CAM. To fulfill this requirement, high accuracy and dimensional stability of a-CAM products are crucial. Based on a recent systematic review, the accuracy of 3D-printed dental models varies widely and can reach a maximum mean error of 579 μm (Etemad-Shahidi et al., 2020). The precision of 3D-printed polymer models is significantly influenced by the digital workflow applied (Mühlemann et al., 2018), the settings of manufacturing parameters (e.g. build direction) (Park et al., 2019) and the type of post-processing and storage (Etemad-Shahidi et al., 2020). It is evident that accuracy of a-CAM products has to achieve the level of s-CAM to be successfully integrated in the digital fabrication workflow.

Further clinical studies on production time applying a digital workflow for the fabrication of implant prostheses should (a) focus on the use of a-CAM technologies, (b) specify the applied a-CAM technology and the parameter settings (e.g. layer thickness) and (c) document the prosthetic and auxiliary components in detail. In addition, the time for post-processing of the a-CAM product (deburring, sintering, cleaning, curing, surface treatments) needs to be recorded. Finally, effectiveness should be reported by the number, type and time of chairside adjustments needed for each prosthesis rather than providing mean time for chairside adjustments.
CONCLUSIONS

The scientific evidence obtained through the present systematic review provides no data on production time, effectiveness and costs in the digital workflow comparing s-CAM and a-CAM. Post-processing of CAM products may substantially contribute to the overall production time. Monolithic CAD-CAM implant crowns may be preferred because the need for chairside/laboratory adjustments was lower compared to veneered implant crowns.

CONFLICT OF INTEREST

The authors report no conflict of interest.

AUTHOR CONTRIBUTION

Sven Mühlemann: Conceptualization (lead); Data curation (lead); Formal analysis (lead); Investigation (lead); Methodology (lead); Validation (lead); Writing-original draft (lead); Writing-review & editing (lead). Jenni Hjerppe: Conceptualization (supporting); Data curation (equal); Formal analysis (equal); Investigation (equal); Methodology (equal); Writing-original draft (supporting); Writing-review & editing (supporting). Christoph H. F. Hämerle: Conceptualization (supporting); Investigation (supporting); Methodology (supporting); Resources (lead); Supervision (supporting). Daniel S. Thoma: Conceptualization (equal); Data curation (supporting); Formal analysis (supporting); Investigation (equal); Methodology (supporting); Writing-original draft (equal); Writing-review & editing (equal).

DATA AVAILABILITY STATEMENT

Data can be provided upon request.

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