UNIVERSITY OF OKLAHOMA GRADUATE COLLEGE

DESIGN AND FABRICATION OF A 3D PRINTED HYBRID IMPLANT FOR TEMPOROMANDIBULAR JOINT REGENERATION

A THESIS SUBMITTED TO THE GRADUATE FACULTY in partial fulfillment of the requirements for the Degree of MASTER OF SCIENCE

By

ALI RASSI Norman, Oklahoma 2021

DESIGN AND FABRICATION OF A 3D PRINTED HYBRID IMPLANT FOR TEMPOROMANDIBULAR JOINT REGENERATION

A THESIS APPROVED FOR THE SCHOOL OF INDUSTRIAL & SYSTEMS ENGINEERING

BY THE COMMITTEE CONSISTING OF

Dr. Pedro Huebner, Chair Dr. Michael Detamore Dr. Shivakumar Raman

© Copyright by ALI RASSI 2021 All Rights Reserved.

Abstract

Patients suffering from temporomandibular joint (TMJ) disorders have difficulties performing trivial activities such as chewing, yawning, and even laughing. In severe cases, the pain is unbearable and clinical care is required. When non- or minimally invasive treatments fall short to resolve the problem, total joint replacements with permanent alloplastic implants are recommended by surgeons. However, recent advances in tissue engineering research have highlighted the potential of using bioengineered implants over their traditional alloplastic counterparts.

In this study, the design and fabrication processes of a patient-fitted hybrid biodegradable TMJ implant are studied, focusing on the mitigation of complications and limitations of conventional implants. The proposed implant design incorporates both bone and cartilage components of the TMJ in a single heterogeneous unit and is affixed to the ramus of the mandible.

By including microporous features in the condylar head of the implant, we seek to increase the rate of cell infiltration via capillary forces in the scaffolding structures and reinforce the bond between the bone and the cartilage regions so as to obtain a faster and more effective regeneration of the targeted tissues. Statistical design of experiments is adopted to identify the best-performing features that are ultimately implemented into the final design of the full-size TMJ implant. Lastly, techniques for incorporating these microarchitectural features and improving the fabrication process are presented and discussed.

Acknowledgments

I would like to thank God for reasons we both know that I will not get into here. Thanks, God.

I want to thank my mom and my aunt too for supporting me and my siblings like we are the only ones ever born to be supported. This thesis is dedicated to you.

I also want to thank Samin and Erfan, my sister and brother. They have not helped with the thesis as much as they could have, but they are hilarious and have said many funny things during the period in which this thesis was written and earned themselves a little something in the acknowledgments section.

I was very lucky to have amazing teammates who made the lab my favorite place on campus. David, Emi, and Boushra thank you for laughing at my jokes and having my back. You guys rock.

I want to thank Dr. Detamore for always being mindful of me and available to help. During my master's program, I had the privilege of knowing many people who were incredibly kind to me and went out of their ways to help me grow. I am also grateful for all the opportunities that Dr. Raman and other faculty and staff from the school of Industrial and Systems Engineering provided for me.

I have to make an un-scientific statement here. Based on my extensive research on the topic, I have concluded that my advisor, Dr. Pedro Huebner, and I are the best duo ever. He has helped me in ways I could not think of and mentored me like no one else would have. A big figure in my life and someone I will look up to. Cannot thank you enough Pedro for believing in me like you did. Your next student is one lucky person.

Table of Contents

1.	I	ntroc	luction	1
1	1.1	Ten	nporomandibular Joint	1
1	1.2	Ten	poromandibular Joint Disorders	2
1	1.3	Clin	ical Relevance	3
	1.	3.1	Current Treatment Options	3
1	1.4	Tiss	ue Engineering for Treatment of TMJ	5
	1.4	4.1	Tissue engineering of the TMJ disc	6
	1.4	4.2	Tissue engineering of the TMJ mandibular condyle	6
	1.4	4.3	Additive Manufacturing	7
1	1.5	The	sis Statement and Outline	8
2.	P	hase	1: Micro-Architecture	.10
2	2.1	Intro	oduction	.10
2	2.2	Mat	erials and Methods	.12
	2.2	2.1	Preparation of the Filament	.12
	2.2	2.2	Optimizing Print Settings	.13
	2.2	2.3	Capillary Action Studies	.14
	2.2	2.4	Preparation of the Hydrogel	.16
	2.2	2.5	Bone-to-Cartilage Integration Studies	.17
2	2.3	Res	ults	.20
	2.3	3.1	Capillary Action Studies Results	.20
	2.3	3.2	Bone-to-Cartilage Integration Studies	.26
2	2.4	Disc	cussions	.32
3.	Р	hase	2: Macro-Architecture	.35
3	3.1	Intro	oduction	.35
3	3.2	Mat	erials and Methods	.37
	3.2	2.1	Design of the Implant	.37
	3.2	2.2	Fabrication of the implant	.39
3	3.3	Res	ults and Discussion	.41

4.	Conclusion	
4.1	1 Summary	43
4.2	2 Contribution	44
4.3	3 Future Works	44
5.	References	45
6.	Appendix	54

List of Figures

FIGURE 2.1 FILAMENT EXTRUDING LINE	13
FIGURE 2.2 CAPILLARY ACTION STUDY TEST SAMPLE	15
FIGURE 2.3 CAPILLARY ACTION STUDY	16
FIGURE 2.4 DELAMINATION RIG	19
FIGURE 2.5 BONE-TO-CARTILAGE INTEGRATION STUDY RIG	20
FIGURE 2.6 UPTAKE RESPONSE	21
FIGURE 2.7 SURFACE TREATMENT EFFECT	26
FIGURE 2.8 PEAK AXIAL FORCE AND STRAIN AT PEAK FORCE	29
FIGURE 2.9 UPTAKE STUDY WELLS	32
FIGURE 3.1 FEATURES OF THE TMJ IMPLANT	39
FIGURE 3.2 IMPLANT PRINT ELEMENTS	41

List of Tables

TABLE 1. BONE-TO-CARTILAGE INTEGRATION STUDY MICRO-ARCHITECTURES. 18
TABLE 2. TESTS OF BETWEEN-SUBJECTS EFFECTS FOR THE DEPENDENT
VARIABLE: UPTAKE
TABLE 3. UNIVARIATE TESTS FOR THE EFFECT OF LAYER HEIGHT
TABLE 4. PAIRWISE COMPARISONS FOR THE EFFECT OF LAYER HEIGHT
TABLE 5. UNIVARIATE TEST FOR THE EFFECT OF LINE DISTANCE 24
TABLE 6. PAIRWISE COMPARISONS FOR THE EFFECT OF LINE DISTANCE 25
TABLE 7. BONE-TO-CARTILAGE INTEGRATION DATA 27
TABLE 8. TESTS OF BETWEEN-SUBJECTS EFFECT FOR DEPENDENT VARIABLE:
PEAK AXIAL FORCE
TABLE 9. TESTS OF BETWEEN-SUBJECTS EFFECT FOR DEPENDENT VARIABLE:
STRAIN AT PEAK
TABLE 10. PAIRWISE COMPARISONS FOR THE EFFECT OF DESIGN ON THE PEAK
AXIAL FORCE
TABLE 11. PAIRWISE COMPARISONS FOR THE EFFECT OF DESIGN ON THE STRAIN
AT PEAK FORCE
TABLE 12. GRAPHICAL VIEW OF THE SURGERY PROCEDURE 36
TABLE 13. GRAPHICAL VIEWS OF THE IMPLANT AND THE CUTTING GUIDE

1. Introduction

1.1 Temporomandibular Joint

We all rely on the function of our joints to perform routine tasks. From pouring water in a kettle for a morning cup of tea to opening the door of our office and driving back home, we take advantage of joints, directly or indirectly, to achieve our goals. We interact with up to a couple of dozens of joints in our day-to-day life. That number, however big it might be, is insignificant when compared to the 360 joints that we use inside our bodies. This number is not constant between all humans, but the range of 300 to 400 is consistent with the literature.[1]

Unlike the joints we utilize in our day-to-day life to help us with our tasks, not all the joints inside our bodies allow for movement. Basically, a joint is the connection between two bones and a common classification of the joints structure in our body, divide them into 3 groups: fibrous joints, cartilaginous joints, and synovial joints. These groups differ in their mobility and range of motion. Some are freely mobile joints (known as diarthrosis), some are slightly mobile (known as amphiarthrosis) and others are totally immobile (known as synarthrosis). While they lie in the general definition of a joint, amphiarthrosis and synarthrosis joints do not provide any functional movement. Synovial joints, on the other hand, allow different types of motions based on their particular structure, thus are of great interest to us as engineers.

Synovial joints can be classified into 3 types based on their range of motion. First, the uniaxial joint, which only allows for movement along one axis, such as rotary move. An example of uniaxial joints would be the rotation of our head around the axis of the neck when we look around. Second, the biaxial joint, which in turn, allows for movement along 2 distinct axes. One might think of the movement of the thumb for this type of joint. The last type is the polyaxial (or multiaxial) joint and, as the name suggests, it allows for movement along 3 or more axes.

The only polyaxial joints are ball-and-socket (or spheroid) joints which are the most mobile joints in the body and can be found in the shoulder and hip. The hinge joints allow for rotation along one axis and can be found in the knee and elbow. Pivot (rotary) joints are also uniaxial and allow for movement along one axis and can be found in the neck. Condylar (or ellipsoid) joints have a unique structure that only allows for movement in 2 axes that are perpendicular to one another. This motion is clearly seen in the wrist, where you can only rotate your hand to either left

and right or up and down. Next, the saddle joint, in which the convex surface of one bone and the concave surface of the other, forms a structure that allows for movement along 2 axes and can be found in the thumb. The last type of joint known as plane joint (or gliding joint), unlike other joints, does not move around an axis and, instead, permits movement on the plane of the articular surface.

One of the less-investigated synovial joints in our body and perhaps the most important one, in that it is responsible for talking, breathing, and chewing food, all of which are essential to the quality of life, is the temporomandibular joint, also referred to as TMJ for short. This load-bearing joint connects the condylar head of the lower jaw (mandible) to the mandibular fossa of the temporal bone, and is the most active joint in our body, with cycles of loading and unloading of up to 2000 times per day.[2]

With the unique structure of the condylar head, TMJ provides movements in the sagittal axis called protrusion/retrusion (anterior/posterior, respectively), in the vertical axis called depression/elevation (lowering/raising, respectively), and in the frontal axis, called lateral deviation. To open the mouth, the TMJ performs a combination of depression and protrusion, and to close the mouth the TMJ movement involves a combination of elevation and retrusion. In lateral movements, left and right TMJs alternate in retrusion and protrusion.[3]

1.2 Temporomandibular Joint Disorders

TMJ disorders or TMDs are referred to a group of disorders in TMJ and the surrounding tissues that in one way or another cause pain or limit the functionality of the joint. These disorders could range from clicking noises during the motion of the mouth to pain in the surrounding tissues of the TMJ, and in some cases, limitation in the range of motion of the TMJ.[4]

There are two main categories for TMJ disorders based on their anatomical origins; masticatory muscle disorders, which are the problems within the muscles surrounding TMJ, and articular disorders, which refer to the problems within the articular surfaces of TMJ. [4] Internal derangement or ID is referred to any interference in smooth joint movement, [5] and several types of IDs are common in TMJ; Disc derangement, disc adherence, disc adhesion, hypermobility, and joint dislocation, to name a few.[6]

The etiology of TMJ disorders is not clear and many studies have hypothesized different factors to have an effect on either initiation or perpetuation of TMDs, some of which have been questioned and in cases, rejected by other studies. [4] Among the most common factors are parafunctional habits such as teeth clenching or teeth griding and it is also possible that non-TMJ disorders such as depression, bad sleeping habits, or rheumatism exacerbate the symptoms of TMD.[7][8][9]

Due to the complex nature of the TMJ and our incomplete understanding of the factors contributing to TMD, preventive actions to control these disorders are very limited, and oftentimes, treatments are necessary after the fact.

1.3 Clinical Relevance

Studies on the prevalence of TMJ-related pains show that anywhere between 25% to 75% of the population have experienced signs of TMD, but only a minority of about 3.6% to 7% have TMD symptoms with such severity that requires them to seek treatment. [2][7] These symptoms have been mostly identifiable in young adults and adults from 20 to 50 years of age, and women have two to three times higher share in the affected population. [10][11]

We should keep in mind that the treatment of TMJ disorders costs about \$4 billion every year in the US alone.[12] A recent study shows that the number of TMJ total joint replacement (TJR) procedures done in the United States, from 412 cases in 2005 has increased to 572 cases in 2014 and is estimated to go above 900 cases in the year 2030, which shows an increased rate of 58%[13]. One of the bigger TMJ solution companies (TMJ Concepts) alone has more than doubled its sales of TMJ TJR devices in the same period, and with the rise of newly FDA-approved devices from different companies, it is expected to have higher demands for such treatments.

1.3.1 Current Treatment Options

The avascular nature of the TMJ does not allow for regeneration and self-healing of the tissues [14] and available treatment methods often target hindering the progress of the disorder [15] and regaining the functionality of the joint, instead of completely healing or regenerating the tissues. Different treatment methods have been advised for people with different levels of TMJ disorder.[16] In many cases, the symptoms of TMD reduce over time, and 85% of people will have no sign in 3 years.[17][18] therefore, non-invasive and minimally invasive treatment options are

recommended for patients in either early stages or with lower severity of TMD, since invasive methods could be harmful and the results are irreversible.[17][7]

Currently, treatment options for TMD include non-invasive, minimally invasive, and invasive, each suggested according to the severity and different stages of the TMD, with the preference given to non-invasive and minimally invasive treatments in the preliminary stages and invasive methods reserved for extreme cases where previous methods could not provide favorable results. [12][19]

Self-care and behavioral interventions are generally considered to be the first step in addressing the TMD and should be employed even if other treatment methods are necessary. Non-invasive therapies are the next step in controlling the TMD and several therapy methods are known to reduce the signs and symptoms. Physical therapies such as lateral jaw movement or resisted jaw opening/closing[20], psychological interventions like cognitive behavior therapy or stress management, dental procedures like occlusal therapy[7][8][9][19], medical interventions such as acetaminophen or anti-depressants, or even hypnosis and acupuncture are said to be effective methods to control TMJ related pains and regain its functionality.[4] However, further investigation is required to directly assess the effectiveness of each of these methods on TMD since, for example, there are studies that conclude and others that reject the improvement in TMD symptoms when physical therapy is used.[21]

An example of a minimally-invasive treatment would be intra-articular injections with local anesthetics or Botox for treatment of chronic bruxism, although it is recommended to only use this method after preservative methods were not sufficient.[17][22] However, there are many studies that show the high effectiveness of this method and due to the simplicity of the procedure, it can be repeated if necessary.[23] This is while another study has associated multiple intra-articular injections with the destruction of articular cartilage.[24] Nonetheless, the necessity of repeated surgery in itself is a sign of a sub-optimal solution for TMD. Other forms of minimally-invasive treatments include arthroscopy and arthrocentesis, both of which have been shown to improve the pain and the range of motion in TMJ patients,[25][26] but there are inconsistencies in the literature about their effectiveness.[27][28]

The last resort in TMJ management is surgery. The American Association of Oral and Maxillofacial Surgeons (AAOMS) has published a list of criteria for eligibility of patients in rare

cases where previous methods have failed to address the pain and functionality of the TMJ that should be assessed before proceeding with TMJ surgery.[29] TMJ surgeries involving disc prosthesis and total joint replacement (TJR) date back to the mid-19th century, but the Proplast-Teflon disc prosthesis once thought to be promising [30] and the majority of experimental methods that surgeons have explored over time only yielded short-time results and often caused more problems in the long term.[31] As of today, there are only 2 FDA-approved manufacturers that produce TMJ replacement devices.

For management of the TMJ disc, surgical methods include arthroplasty and disc repositioning,[32] eminectomy,[33] discoplasty,[34] and discectomy.[35] None of these treatments provide permanent results and follow-up therapies or surgeries are often needed.[36] For that reason, the development of methods that assure long-term results are imperative. It is worth mentioning, however, that according to a study on more than 2000 TMD patients, only 2.5% had to receive surgery and it should only be done when non- or minimally invasive methods have not been effective.[37]

1.4 Tissue Engineering for Treatment of TMJ

As previously mentioned, many treatment approaches have emerged over the past several decades to mitigate the symptoms of TMD, hinder its progress, and regain the functionality of the TMJ, some of which are still in use today. Unfortunately, none of the currently available methods targets complete healing or restoration of the affected joint and its surrounding tissues, and as a consequence, patients often need follow-up surgeries.[36] Besides, the few treatment options that are known to be successful are only suitable for a portion of the affected population who can pass a long checklist of eligibility criteria.[29] Thus, it is clear that there is a need for new treatment methods that could address these gaps and produce better and more desirable results for a wider population. In recent years, tissue engineering has been considered a promising solution for the regeneration of the TMJ.

There are indications for the use of tissue engineering in the reconstruction of the TMJ components, such as the TMJ disc and the mandibular condyle, and several studies have explored these areas. The following is a brief overview of the past and current endeavors in this domain.

1.4.1 Tissue engineering of the TMJ disc

From the literature, patients with trauma or internal derangement (ID) are the main groups who could benefit from bioengineered discs. As mentioned in <u>chapter 1.3.1</u> a standard for treatment of ID where the disc cannot be repositioned or repaired is discectomy, a surgery where the disc is completely removed. Even though this is the last resort, the results of discectomy are far from encouraging [38] and TE is considered to be a solution to replace the damaged disc and potentially avoid the discectomy. [39]

Studies on tissue engineering of the TMJ disc have tried to recreate/restore the disc through scaffold-based and scaffold-free approaches. [12] TE in these studies aims at making a scaffold that 1) mimics the anatomy of the tissue 2) grants an equal or comparable rate of degradation for the scaffold and rate of synthesis for the tissue. Many synthetic materials such as Polylactic Acid (PLA) [40], Poly(L-lactide) (PLLA), [41] polyglycolic acid (PGA), [42], and Polyethylene Glycol Diacrylate (PEGDA) [43], as well as natural scaffolds like extracellular matrix[43][44], have been examined to find an appropriate configuration.

In cases where the scaffold degradation has significantly surpassed the synthesis of the tissue, efforts have been made to either slow down the degradation rate via different material selection, or speed up the growth of native tissues with growth factors such as insulin-like growth factor, basic fibroblast growth factor, and transforming growth factor. [42][41]

More closely related to the scope of this thesis, one recent study has utilized 3D printing to closely mimic the shape and tissue alignment of TMJ disc using PCL, and with the help of multiple growth factors embedded in different areas of the disc, they could achieve a tissue-engineered disc with similar anatomy to that of the native disc, and the right configuration of fibrous/fibrocartilaginous matrix in the corresponding areas.[45]

1.4.2 Tissue engineering of the TMJ mandibular condyle

Generally, 4 groups of patients are potential candidates for tissue-engineered mandibular condyle and ramus. The first group is patients with severe condylar trauma where the condyle is not salvageable. In these cases, biological reconstructions and/or TMJ implants are advised for the reproduction of the anatomical form.[46][47] The second group is adolescents and young adults who are still in the age of growth. In these cases, the use of alloplastic implants requires follow-

up surgeries to keep up with the growth of the patient. To resolve that issue, experiments have been done with autografts with hopes of growing with the patient, [48][49][47] but there are instances where the un-equal rate of growth between the native tissue and the autograft has caused failure. [50][51][52][53] The third group is patients with condylar hyperplasia, which is a condition where excessive cell proliferation results in an increased amount of organic tissues. Bioengineered implants could potentially replace the diseased tissues and hinder the progress of hyperplasia. [46] The last group is those with metal hypersensitivity where the use of bioengineered condylar implants can prevent the adverse reaction of the patient to alloplastic implants. [54]

A thorough review of past efforts on tissue engineering of TMJ condyle by Detamore *et al.* [55] and a newer study by Acri *et al.* [56] explore the recent advancements of TE in this realm. Focus on this area should be on engineering the condyle in a way that bone and cartilage regeneration are both achieved while mechanical requirements of the TMJ environment are met. Mechanical properties are usually controlled through design parameters of the condyle scaffold such as pore size and porosity, and biochemical characteristics of such implants are commonly modified via material selection and post-processing procedures to ensure the biocompatibility, scaffold degradation rate, and tissue regeneration rate.

As for the tissue engineering of the condyle, both synthetic and natural materials have been explored for the tissue engineering of the mandibular condyle. Among synthetics, PLA,[57] PGA,[58] PLGA,[59] PCL,[60] and HA[61] and among natural materials, decellularized bone matrix,[62] coral,[63] collagen,[64] and chitosan[65] are the most prominent scaffolding materials

1.4.3 Additive Manufacturing

As opposed to subtractive manufacturing technologies where the final part is obtained through subtraction from a bigger volume of raw material, Additive manufacturing (AM) is a process in which a part is fabricated by stacking up layers of materials on top of each other. This process, also known as 3D printing, allows the fabrication of very complex structures that are otherwise impossible with traditional methods.[66] The ease of use and flexibility of AM has made it interesting to many sectors, and medicine and healthcare are among the fields where AM popularity is increasing at a rapid rate.[67] The main applications of AM in the biomedical sector include:

- fabrication of tools, instruments and equipment [68]
- physical models for visualization, preoperative planning,[69] testing, and education
- fabrication of customized implants[70]
- biostructures for scaffolds and tissue engineering[60][45]

At its core, FFF (Fused Filament Fabrication) consists of a set of carriers controlled by stepper motors that move in either cartesian or polar coordinates and position the print head on a specific point on top of a print bed. Another stepper motor is in the print head where an extruder is heated up to the melting temperature of the material. This material is usually fed to the printer as a strand called filament. The print head moves in a defined path to deposit material and then moves away from the print bed and begins to print another layer on top of the previous layer. This process repeats until a complete model is printed. This configuration (in cartesian 3D printers) requires 4 stepper motors to produce the desired shape; 3 to control the position of the head, and one to extrude material. The defined path is generated by a software (known as a slicer) that slices the desired model into layers and controls the amount of extrusion at any given point of the print.

1.5 Thesis Statement and Outline

Regenerative medicine and additive manufacturing have emerged as promising techniques to address problems that conventional treatment options fail to resolve. Using the principles of regenerative medicine and utilizing the capabilities of 3D printing, in this paper we present a hybrid implant that aims at the regeneration of the temporomandibular joint.

In phase 1, through design of experiments, I will prove that by modifying the design of micro-architectures we can change their response to different factors. In my first experiment, I will show that by changing the print parameters and surface features of a scaffold, we can induce different capillary forces and in turn, adjust its capacity to transport solutions such as bone marrow. In my second experiment, I will study the effect of micro-architectures on the integration of the scaffold and the hydrogel. The results of this experiment will then be used to create stronger bonds between the bone and the cartilage elements of my hybrid implant. From the translational point of view, phase 1 is of high importance because the micro-architectures have the potential to be used in other implants regardless of their shape. This phase also includes protocols for preparing the materials used in my studies.

In phase two, I will present a framework for the design and fabrication of a full-size hybrid TMJ implant. I will then use the findings from my experiments to incorporate selected microarchitectures from phase 1 in the full-size implant that could improve its performance. Additional features will be added to the implant to facilitate the surgeon's job during the surgery. The last chapter includes a summary of the results, discussions, and contributions presented in this thesis and an outline of opportunities for future work.

2. Phase 1: Micro-Architecture

2.1 Introduction

In every manufacturing process, the material used is chosen to satisfy the requirements of a certain purpose and 3D printing is no different. Many 3D printing materials have been developed to address different needs [71] and each has its own set of parameters under which the manufacturability of the material is optimized. These parameters include the temperature at which the material is extruded, the speed of extrusion, and the range of nozzle sizes the materials could be extruded from. [72][73] and these parameters are usually provided by the material manufacturer or could be found online for the more common materials. Sometimes, the commonly available materials are not appropriate for certain purposes and custom-made materials must be developed to meet specific requirements such as degradation rate or tensile strength.[74]

Knowing the underlying principles of FFF 3D printing, it is important to note that the generated path by the slicer is independent of the material used for the print. Each material has its own characteristics and they should be taken into consideration when slicing a model as they greatly affect the success rate and the quality of the finished part.[75] If known, these parameters could be entered into the slicer package to generate a tool path for that specific material. But in case your material is custom-made and there is no data available online, then you could determine those attributes through a series of experiments. Finding the print settings for optimal manufacturability will be the topic of section 2.2.2.

Generally, when one speaks of "quality of print", it either refers to the surface quality of the part, meaning its appearance is satisfactory for the intended purposes, usually for ornamental parts, or it refers to the mechanical quality of the part, meaning how it performs under a set of mechanical tests, usually in industrial parts. Surface quality is more subjective and depends on a person's judgment, while mechanical performance is more objective and could be assessed to meet your requirements. Nonetheless, methods exist to find the optimal print setting for each material in both appearance and performance quality. Although the performance of a part is heavily determined by the material used, design modifications could be used to achieve better performances [76][77] and that will be discussed in <u>section 2.4</u>.

Extruding filament is a straightforward process and mostly requires a filament extruder and the material you want to have in filament form. The filament extruder has a heated area where the material is pushed through and the molten material comes out of a nozzle at the end of the machine. The material is usually fed to the machine either in pellet, powder, or shredded form through a hopper attached to the heated area.

The precision of the filament is vital to the surface quality of the final print. One of the parameters entered into the slicer is the diameter of the filament. The slicer then calculates the volume of material needed to create the deposited line with a certain height and width and runs the stepper motor to extrude material based on that calculation. This is assuming that the diameter of the filament is constant when in reality there is always variability in the diameter. So, for example, when the diameter of the filament reduces at a certain point of the filament, this results in a deposited line thinner than the ones before and after it and could be easily seen on the surface of the print. Although simple, extruding a good filament requires several measures to minimize the variability in the filament diameter and we have incorporated a few of them in our filament extruding line. The prepared materials were used in the fabrication of test samples for the two experiments that are included in this phase; the capillary action studies, and the bone-to-cartilage integration studies.

Including micro-porous structures in the scaffold is known to increase the volume and rate of bone regeneration by inducing capillary forces.[78] In the capillary action studies, I seek to determine whether it is possible to increase the rate and capacity of uptake in the implant by changing the design of micro-architectures in the scaffold. If successful, the results of this experiment could be used to incorporate porosity with the right parameters in parts of our implants where bone regeneration is desired.

As mentioned in <u>section 1.1</u>, the opening of the mouth involves a rotational and a translational movement of the condylar head, during which the condyle slides against the TMJ disc. The cartilage layer that sits on top of the condylar head allows these movements to be smooth. In the bone-to-cartilage integration studies, I will experiment with micro-architectures that could integrate and sustain the hydrogel element on top of the scaffold during TMJ movements. This study is designed to find the design that maximizes the forces it takes for the hydrogel to be sheared/delaminated off of the implant scaffold.

2.2 Materials and Methods

2.2.1 Preparation of the Filament

Polycaprolactone (PCL) filament with 20% weight hydroxyapatite powder (HAp) was made using a filament extruder (Noztek Touch). PCL (Polysciences, Inc.) and HAp (Aldrich) were mixed with the desired ratio and ball-milled for 19 hours. The powder was stored in a desiccator prior to extruding to evaporate any moisture in the mix, as it noticeably reduces the quality of print.

The filament extruder heats the material in two phases using two separate heating elements. The speed of extruding the filament and the temperature for both heating elements are controlled via the touchscreen panel on the machine. The first and the second heating elements were set to 80 °C and 140 °C, respectively. The speed of the extruding motor was set to 30 rpm. Figure 2.1 shows the units that make up the filament extruding line.

The extruded filament was then directed towards a U-shaped aluminum profile of approximately 1m lengths, filled with deionized water that is placed in an ice bath. After cooling down and solidifying, the filament was fed to the puller and the speed of pulling was adjusted to achieve the correct diameter. The filament was made in both 1.75 mm and 2.85 mm diameters for use with different 3D printers.

A similar gearbox to that of a direct drive 3D printer extruder was used in place of a puller to control the speed with which the filament is being pulled. The stepper motor in the puller unit is connected to an Arduino Uno and the speed is controlled via a potentiometer or the serial monitor. The extruded filament was then coiled around a filament spool using a filament winder (Noztek Filament Winder 1.0).



Figure 2.1 Filament Extruding Line

No more than 60 grams of powder was extruded in one run to maximize the quality of filament and prevent the excessive build-up of residues on the walls of the filament extruder. The filament extruder was thoroughly cleaned after each run so that there is no residue left from previous runs.

2.2.2 Optimizing Print Settings

Since 80% of our material consists of PCL, it was the logical choice to start the setting optimization from the available data on the PCL print settings. A cylinder of diameter 10 mm with an interconnected porous network was chosen as the test sample to determine the effect of different attributes on the quality of the printed part.

The melting temperature of PCL is anywhere between 60 °C to 100 °C in the literature [79][73], so the starting point for our print temperature was set to 60 °C. Using a post-processing plugin in Ultimaker Cura called Tweak At Z, the temperature was gradually increased with height, spanning a range of temperatures from 60 °C to 160 °C.

It was noticed that PCL takes much longer to solidify since the temperature drop it experiences once extruded is much lower than other commonly used materials and this necessitates the use of active cooling fans. After turning on the cooling fans it was noticed that the printed part only solidifies where it is closer to the fans. Custom channels were designed to cool down the printed parts in a more uniform way. This addition visibly improved the quality of the printed parts.

After determining a feasible print temperature, print speed was decreased from 30 mm/s to 1 mm/s, and the flow rate was increased from 80% to 120%, following the same method. Print bed temperature was decreased from 80 °C to 30 °C for separate test samples since it only affects the first layer of the print. Other settings such as retraction, fan speed, and minimum layer time were modified after a satisfactory print was achieved to optimize the quality.

The material was ultimately printed with the following settings:

- print temperature: 135 °C
- bed temperature: 38 °C
- print speed:
 - shell speed: 10 mm/s
 - support speed: 10 mm/s
 - infill speed: 2.5 mm/s
- fan speed: 100%
- flow rate: calculated based on the diameter of the filament

2.2.3 Capillary Action Studies

Test samples were made to determine the response of capillary actions to different microarchitectures. Porous cubes of size $12x12x12 \text{ mm}^3$ with 3 different pore sizes (0.6 mm, 0.8 mm, 1.0 mm line distance) and 3 different layer heights (0.12 mm, 0.18 mm, 0.3 mm) were used for the capillary action studies. Each group had 3 samples (n=3) and all samples were cleaned with 70% ethanol and then air-dried to remove any grease or dirt before the experiments were done.



Figure 2.2 Capillary action study test sample.

A, *B*) *Test samples were printed with a zigzag pattern to form an interconnected porous network. C*) *Cross-section of a test sample. The values for line distance, line width, and layer height are entered into the slicer.*

Each sample was characterized individually on a 40x magnification microscope. 3 random areas were chosen to take measurements of the pore size and line width of each sample.

Red food coloring was added to 1 liter of PBS and the colored solution was used to identify the height of uptake across all test samples. Wells of 1 ml were filled with 500 μ l of the uptake solution using a pipette. The wells were filled halfway to account for the volume of the test samples when inserted to prevent overflowing. The dry test samples were weighed, placed into the solution pool for 5 minutes, and then weighed again. The difference between the dry weight (before) and the wet weight (after) is the amount of uptake induced by each sample. Pictures were taken at the start and end of the 5 minutes. It was recorded if the solution has reached the top of the sample, as it means that the uptake amount was limited by the size of the well and the test sample.



Figure 2.3 capillary action study

Treated test samples with 0.8 mm line distance and 0.12 mm layer height after 5 minutes of soaking in the uptake solution. Notice how in the middle sample the solution approaches the top of the cube but does not reach it.

Test samples were then washed, air-dried, and etched with Sodium Hydroxide (NaOH, VWR chemicals BDH[®]) to measure the effect of surface treatment on the capillary forces. NaOH pellets were mixed with DI water to reach a concentration of 5mol/liter. Test samples were soaked in the NaOH solution for 30 minutes and then washed with DI water and air-dried 3 times. The uptake study in the same procedure as mentioned above was repeated to measure the amount uptake for the treated test samples. Figure 2.3 shows three samples after 5 minutes of soaking in the uptake solution.

2.2.4 Preparation of the Hydrogel

GelMA Lyophilizate (CELLINK) at 5% (w/v) was prepared in the following procedure. Lithium phenyl-2,4,6-trimethyl-benzoyl phosphinate (LAP, CELLINK) was dissolved in Phosphate-Buffered Saline (PBS) at 0.6475 mg/ml and GelMA was added to this solution at 50 mg/ml. Polyethylene Glycol Diacrylate (PEG-DA average M_n 4,000, Aldrich) was added to the solution until it reached 10% (w/v) of the mix. The final mix was placed in a bead bath at 37 $^{\circ}$ C and then vortexed until the GelMA was dissolved and the mix was homogeneous. The mix was cooled down to room temperature (25 $^{\circ}$ C) and stored away from light.

2.2.5 Bone-to-Cartilage Integration Studies

Test samples with different geometries were made to assess the effect of different microarchitectures on bone-to-cartilage integration. These samples consisted of the micro-architecture most successful in the capillary action studies with the addition of features that could potentially reinforce the bond between the hydrogel and the scaffold. Different designs included:

- zigzag-to-ribs:
 - \circ from 0.6 mm zigzag on the bottom half to 0.6 mm ribs on the top half
 - \circ from 0.6 mm zigzag on the bottom half to 0.8 mm ribs on the top half
 - 0 from 0.6 mm zigzag on the bottom half to 1.0 mm ribs on the top half
- rectangular slot:
 - \circ shallow (3x8x3 mm³)
 - \circ deep (3x8x6 mm³)
- cross slot:
 - 0 horizontal
 - 0 vertical
- anchorage:

• 2 slots (3x3x3 mm³) connected through a channel (3x8x3 mm³) A schematic representation of these designs is shown in table 1.





Prior to conducting the study, samples were etched and washed as established in <u>chapter</u> 2.2.4. The hydrogel prepared above (in <u>chapter 3.2.2</u>) was manually syringed inside the test samples that had slots/anchors. These samples were then placed on the build plate of the BioAssemblyBot (BAB, Advanced Solutions Life Sciences) and were registered with a 3D representation of the same sample inside the BAB's dedicated 3D modeling software Tissue Structure Information Modeling (TSIM). A rectangular box of size 10x10x2 mm³ was printed on top of all samples with the hydrogel. Test samples were crosslinked for 10 minutes before the integration study.

A testing rig was custom-designed in Fusion 360 (Autodesk) for the Discovery HR-2 rheometer (TA Instruments) to conduct studies on the bond between the hydrogel and different micro-architectures (figure 2.4). This attachment consists of a shear head that attaches to the tool holder of the rheometer, a base that sits on the Peltier plate of the rheometer, a panel that slides on top of the base to change the position of the test sample, a guiding screw inside the panel, and a grip that is mounted on the guiding screw. This grip is tightened to the sliding panel at the correct position to hold the test sample in place for the integration study.



Figure 2.4 Delamination rig

The shear head was lowered until it reached the top of the hydrogel. The test sample was positioned so that the shear head was aligned with the interface of the bone-to-cartilage region. An axial force procedure was performed on the rheometer and the data for the forces it takes to delaminate the hydrogel off of the test samples was recorded.



Figure 2.5 Bone-to-cartilage integration study rig

A) The shear head is lowered until it reaches the hydrogel. The test sample is positioned such that the shear head only contacts the hydrogel. After contact, the hydrogel is either B) delaminated/sheared off, or C) morphed until it catastrophically fails.

2.3 Results

2.3.1 Capillary Action Studies Results

The weight of uptake in each sample was the dependent variable and was analyzed based on three parameters of layer height, line distance, and surface treatment. The geometries with 0.6 mm line distance and 0.12 mm layer height (Mean = 253 mg, SD = 211) and 0.8 mm line distance and 0.18 mm layer height (Mean = 50 mg, SD = 28) scored the highest and the lowest, respectively, out of all 9 different micro-architectures. The descriptive statistics of this experiment are included



in the appendix. The effect of different factors on the uptake can be seen in figure 2.6. These charts are constructed based on the average uptake amount across all test samples.

Figure 2.6 uptake response

The response of uptake to factors line distance, layer height, and surface treatment. Error bars show 1 standard deviation for each group. Dashed lines indicated significant difference between groups (α =0.05)

Based on the three-way ANOVA analysis of these three factors with 95% level of confidence ($\alpha = 0.05$), we can conclude that layer height (p < 0.00) and surface treatment (p < 0.00) have a significant effect on the amount of uptake induced by each sample, while line distance was only borderline significant (p = 0.0504). But to interpret these p-values, we first have to determine the significance of their interactions.

Table 2. Test	ts of between-	subiects effe	cts for the de	evendent va	riable: Uptake

Dependent Variable: Uptake								
Type III Sum								
Source	of Squares	df	Mean Square	F	Sig.			
Corrected Model	541758.139 ^a	17	31868.126	9.887	.000			
Intercept	725904.978	1	725904.978	225.204	.000			
LayerHeight	133969.425	2	66984.712	20.781	.000			
Etch	206894.560	1	206894.560	64.187	.000			
LineDistance	20945.753	2	10472.876	3.249	.050			
LayerHeight * Etch	79590.611	2	39795.306	12.346	.000			
Etch * LineDistance	36261.024	2	18130.512	5.625	.007			
LayerHeight * LineDistance	32841.053	4	8210.263	2.547	.056			
LayerHeight * Etch * LineDistance	31255.713	4	7813.928	2.424	.066			
Error	116039.373	36	3223.316					
Total	1383702.490	54						
Corrected Total	657797.512	53						

a. R Squared = .824 (Adjusted R Squared = .740). Blue cells indicate significant effect.

From table 2, the interaction between the layer height and the surface treatment seems to be significant (p < 0.00), so to conclude that the main effect of each of these factors is truly significant, we conduct a simple main effect analysis (also called univariate test) to find the effect of layer height within treated and un-treated groups.

Table 3. Univariate tests for the effect of layer height

Dependent Variable: Uptake

H	Etch	Sum of Squares	df	Mean Square	F	Sig.
no	Contrast	5359.642	2	2679.821	.831	.444
	Error	116039.373	36	3223.316		
yes	Contrast	208200.394	2	104100.197	32.296	.000
	Error	116039.373	36	3223.316		

Each F tests the simple effects of LayerHeight within each level combination of the other effects shown. These tests are based on the linearly independent pairwise comparisons among the estimated marginal means. Blue cells indicate a significant effect.

Table 4. Pairwise comparisons for the effect of layer height

						97.5% Confidence	
			Mean			Interval for	Difference ^b
	(I)	(J)	Difference	Std.		Lower	Upper
Etch	LayerHeight	LayerHeight	(I-J)	Error	Sig. ^b	Bound	Bound
no	.12	.18	29.589	26.764	.276	-33.013	92.191
		.30	589	26.764	.983	-63.191	62.013
	.18	.12	-29.589	26.764	.276	-92.191	33.013
		.30	-30.178	26.764	.267	-92.780	32.424
	.30	.12	.589	26.764	.983	-62.013	63.191
		.18	30.178	26.764	.267	-32.424	92.780
yes	.12	.18	214.422*	26.764	.000	151.820	277.024
		.30	121.956*	26.764	.000	59.354	184.557
	.18	.12	-214.422*	26.764	.000	-277.024	-151.820
		.30	-92.467*	26.764	.001	-155.068	-29.865
	.30	.12	-121.956*	26.764	.000	-184.557	-59.354
		.18	92.467*	26.764	.001	29.865	155.068

Dependent Variable: Uptake

Based on estimated marginal means. Blue cells indicate a significant effect.

*. The mean difference is significant at the .025 level.

b. Adjustment for multiple comparisons: Least Significant Difference (equivalent to no adjustments).

Table 3 shows that the effect of layer height when the samples are not etched is insignificant on the amount of uptake (p = 0.44). This is while in the treated group, the layer height seems to be having a significant effect (p < 0.00). To interpret this significance, we have to look at the pairwise comparisons of different levels of layer height when they are etched. From table 4 it can be seen that when treated, layer height has a significant effect on the uptake amount in all three levels.

Following the same logic as above this time for the interaction between the line distance and the surface treatment, we can conclude that the effect of line distance on the amount of uptake is only significant when the samples are etched. Within the treated group, there was a significant difference only between 0.6mm-0.8 mm line distance and 0.6 mm-1.0 mm line distance groups.

Dependent Variable: Uptake							
		Sum of					
F	Etch	Squares	df	Mean Square	F	Sig.	
no	Contrast	1107.742	2	553.871	.172	.843	
	Error	116039.373	36	3223.316			
yes	Contrast	56099.034	2	28049.517	8.702	.001	
	Error	116039.373	36	3223.316			

Each F tests the simple effects of LineDistance within each level combination of the other effects shown. These tests are based on the linearly independent pairwise comparisons among the estimated marginal means. Blue cells indicate a significant effect.

Table 6. Pairwis	e comparisons	for the	effect	of line	distance
------------------	---------------	---------	--------	---------	----------

		1		Ĩ		97.5% Confidence	
			Mean			Interval for	Difference ^b
	(I)	(J)	Difference	Std.		Lower	Upper
Etch	LineDistance	LineDistance	(I-J)	Error	Sig. ^b	Bound	Bound
no	.6	.8	-15.489	26.764	.566	-78.091	47.113
		1.0	-9.911	26.764	.713	-72.513	52.691
	.8	.6	15.489	26.764	.566	-47.113	78.091
		1.0	5.578	26.764	.836	-57.024	68.180
	1.0	.6	9.911	26.764	.713	-52.691	72.513
		.8	-5.578	26.764	.836	-68.180	57.024
yes	.6	.8	102.844*	26.764	.000	40.243	165.446
		1.0	89.067*	26.764	.002	26.465	151.668
	.8	.6	-102.844*	26.764	.000	-165.446	-40.243
		1.0	-13.778	26.764	.610	-76.380	48.824
	1.0	.6	-89.067*	26.764	.002	-151.668	-26.465
		.8	13.778	26.764	.610	-48.824	76.380

Dependent Variable: Uptake

Based on estimated marginal means. Blue cells indicate a significant effect. The mean difference is significant at the .025 level.

b. Adjustment for multiple comparisons: Least Significant Difference (equivalent to no adjustments).

Although the effect of surface treatment was not uniform across different geometries, all test samples absorbed more uptake solution after surface treatment, and in the geometry with the biggest improvement, the amount of uptake has increased by more than 7-fold. A schematic representation of the treatment effect is provided in figure 2.7.



Treatment Effect On Different Geometries

Figure 2.7 surface treatment effect

effect of surface treatment on the amount of uptake across different micro-architectures. A) 0.12 mm layer height group. B) 0.18 mm layer height group. C) 0.30 mm layer height group. Dashed lines indicated significant difference between groups $(\alpha=0.05)$.

2.3.2 Bone-to-Cartilage Integration Studies

From the rheology data, peak axial force and strain at peak force were extracted for each sample. The rheology data of axial force vs. strain is shown in table 7. The yellow line indicates the average of 3 replicates within each design. Groups with the deep slot feature (Mean = 1.99 N, SD = 0.54) and the 0.6 mm ribs (Mean = 0.90 N, SD = 0.24) scored the highest and the lowest, respectively. The complete descriptive statistics for the delamination study are included in the appendix.

The graphs in table 7 represent the axial force vs. strain %. The data was adjusted to only show the portion of the experiment where the shear head was in contact with the hydrogel and before the hydrogel was either delaminated or catastrophically failed. Before contact and after

failure the axial force was dropped to zero and the recorded data did not have any valuable information.



Table 7. bone-to-cartilage integration data





Figure 2.8 Peak axial force and strain at peak force

Peak axial force and strain at peak force across all replicates of all micro-architectures. Note how the average of peak axial force and strain at peak force does not differ much within the ribs group and between shallow slot, horizontal cross, vertical cross, and anchored micro-architectures. Dashed lines indicated significant difference between groups (α =0.05).

The force required to delaminate printed hydrogel on top of different micro-architectures as well as displacement of the gel was recorded. These data were analyzed in separate one-way ANOVAs with 95% level of confidence ($\alpha = 0.05$) to determine the significance of each design on the strength of bone-to-cartilage integration.

Table 8. tests of between-subjects effect for dependent variable: Peak Axial Force

lests of Between-Subjects Effects							
	Type III						
	Sum of		Mean				
Source	Squares	df	Square	F	Sig.		
Corrected	3.054a	7	.436	3.147	.027		
Model							
Intercept	46.872	1	46.872	338.10	.000		
				1			
design	3.054	7	.436	3.147	.027		
Error	2.218	16	.139				
Total	52.144	24					
Corrected	5.272	23					
Total							

of Dotrucon **—** .

a. R Squared = .579 (Adjusted R Squared = .395). Blue cells indicate significant effect.

Table 9. tests of between-subjects effect for dependent variable: strain at peak

Tests of Between-Subjects Effects							
	Type III						
	Sum of		Mean				
Source	Squares	df	Square	F	Sig.		
Corrected	1115.439 ^a	7	159.348	8.229	.000		
Model							
Intercept	6309.568	1	6309.568	325.835	.000		
design	1115.439	7	159.348	8.229	.000		
Error	309.829	16	19.364				
Total	7734.836	24					
Corrected	1425.268	23					
Total							

a. R Squared = .783 (Adjusted R Squared = .688). Blue cells indicate significant effect.

Tables 8 and 9 suggest that the design of micro-architectures has a significant effect on the peak axial force and strain at peak force. However, this significance might have arisen from the difference between certain groups and not all groups. To determine the effect of each design, a pairwise comparison test was conducted for both peak axial force and strain at peak force.

Pairwise Comparison: Peak Axial Force	0.6 mm ribs	0.8 mm ribs	1.0 mm ribs	Shallow Slot	Deep Slot	Horizontal Cross	Vertical Cross	Anchorage
0.6 mm ribs	-	0.759	0.734	0.044*	0.002*	0.036*	0.051**	0.056
0.8mm ribs	0.759	-	0.974	0.079	0.005*	0.066	0.091	0.100
1.0 mm ribs	0.734	0.974	-	0.084	0.005*	0.070	0.096	0.106
Shallow Slot	0.044*	0.079	0.084	-	0.183	0.923	0.939	0.896
Deep Slot	0.002*	0.005*	0.005*	0.183	-	0.214	0.161	0.147
Horizontal Cross	0.036*	0.066	0.070	0.923	0.214	-	0.863	0.821
Vertical Cross	0.051**	0.091	0.096	0.939	0.161	0.863	-	0.957
Anchorage	0.056	0.100	0.106	0.896	0.147	0.821	0.957	-

Table 10. pairwise comparisons for the effect of design on the Peak Axial Force

Table 11. pairwise comparisons for the effect of design on the Strain at Peak Force

Pairwise Comparison: Strain at Peak force	0.6 mm ribs	0.8 mm ribs	1.0 mm ribs	Shallow Slot	Deep Slot	Horizont al Cross	Vertical Cross	Anchora ge
0.6 mm ribs	-	0.831	0.550	0.022*	0.000*	0.004*	0.003**	0.014*
0.8 mm ribs	0.831	-	0.699	0.033*	0.000*	0.006*	0.004*	0.022*
1.0 mm ribs	0.550	0.699	-	0.071	0.000*	0.014*	0.009*	0.048
Shallow Slot	0.022*	0.033*	0.071	-	0.004*	0.415	0.318	0.841
Deep Slot	0.000*	0.000*	0.000*	0.004*	-	0.021*	0.031*	0.006*
Horizontal Cross	0.004*	0.006*	0.014*	0.415	0.021*	-	0.848	0.537
Vertical Cross	0.003**	0.004*	0.009*	0.318	0.031*	0.848	-	0.421
Anchorage	0.014*	0.022*	0.048	0.841	0.006*	0.537	0.421	-

From tables 10 and 11 it can be interpreted that the significance of design on the peak force and strain at peak force is mostly due to the difference between the ribs group and the rest of the micro-architectures. Within the ribs group, none of the designs had a significant effect on the peak force compared to one another, and the same could be said for the rest of the designs.

2.4 Discussions

It is evident from the capillary action study data that different geometries in fact induce different capillary forces. This means we can tailor our architecture in order to expedite the rate of bone marrow absorption and increase the height it can reach inside our implant. Utilizing these capillary forces we can ensure that the bone marrow reaches the hydrogel region on top of the implant.

Pictures taken during this study show that all the wells belonging to samples with 0.12 mm layer height and 0.6 mm line distance were empty of the uptake solution after the experiment (figure 2.9), indicating that the test samples absorbed all the solution.



Figure 2.9 Uptake study wells

Empty wells after the uptake study belonging to samples with 0.6 mm line distance and 0.12 mm layer height

Since some of the samples absorbed all of the solutions from their wells, the data might have been different if the samples were placed in bigger wells and with more solutions. Regardless, because all those samples belonged to a certain geometry (0.12 mm layer height and 0.6 mm line distance), this does not affect our judgment on choosing the geometry with the highest capillary forces. Nonetheless, it would be interesting to repeat the same experiment with the same geometry but on bigger scales (bigger wells and bigger test samples) to find the actual capacity of this architecture.

Based on the results of these capillary action studies, the test sample with 0.6 mm line distance and 0.12 mm layer height is the geometry that induced the highest amount of uptake and therefore should be used in the implant design.

The height of test samples at 12 mm were taller than the average height of condylar heads at 8.25 mm (calculated from the 6 patients these implants were designed for), thus if an architecture has succeeded in transporting the solution to the top of the test sample, it is expected for the bone marrow to reach the hydrogel region given the same architecture is used in the implant.

It is also important to consider the difference between our uptake solution and bone marrow. While the study determines the highest capillary forces across different geometries, it is possible that the results would be different with bone marrow.

Although each test sample has been placed in the uptake solution for 5 minutes, most of the absorption has taken place instantly or within seconds of contact with the solution.

After determining the optimal micro-architecture for capillary actions, an experiment was conducted to find the micro-architecture that creates the strongest bond between the hydrogel and the scaffold. The result of this experiment could then be used in the interface of cartilage and bone to integrate the two together.

It was noticed that simply printing a hydrogel on top of the test samples does not create a strong bond between the two phases and that the gel tends to float on the surface of the scaffold. To resolve this weak integration, different designs were studied. In all proposed designs, the body of each test sample consisted of the micro-architecture with the highest capillary forces found in chapter 2.2.3 to ensure that bone-to-cartilage integration does not negatively affect the capacity of the scaffold for bone marrow uptake and features were added to these samples only where they were needed.

In the ribs group, the upper half was designed with the intention of allowing the hydrogel to infiltrate deeper into the structure compared to the zigzag design while the group with the rectangular slot assessed the response of axial force to the thickness of the hydrogel. The rationale behind the design of the cross slot group was to determine the effect of step-like structures inside the test samples. This effect especially became interesting after the shallow slot test samples were studied and two out of three samples failed with the hydrogel delaminating from the scaffold as

one piece. This is while in the deep slot, cross slot group (vertical and horizontal), and anchorage micro-architectures, the hydrogel was not delaminated, rather failed catastrophically. It can be seen in figure 2.7 that the sample with the shallow slot that did not delaminate as one piece (replicate 2) has a higher peak axial force compared to the other two. This means that trapping the hydrogel on the surface of the scaffold could increase the peak axial force.

Another consideration was the strain at which the hydrogel was delaminated or failed. This is important because even if a micro-architecture out-performs other designs at peak axial force, but ceases to hold the hydrogel intact at a very low strain percentage, it will ultimately fail once implanted. The optimal configuration will withstand both the force and strain that the native condylar head experiences in the TMJ.

In the ribs group, due to weak integration, the hydrogel was delaminated in all test samples and the experiment was finished at a very low strain percentage. The hydrogel on other microarchitectures did not delaminate and endured a higher strain before failure.

The bone-to-cartilage integration study suggests that the design with the deep slot was the most successful micro-architecture in both the peak axial force and strain at peak force factors and with the implementation of this micro-architecture on the interface of the hydrogel and the scaffold, we can hope to improve the performance of our implant.

3. Phase 2: Macro-Architecture

3.1 Introduction

In phase one we showed that by using certain micro-architectures we can improve the performance of our test samples. In phase two we will utilize our findings from phase one to design a full-size TMJ implant.

In this paper our aim is to design and fabricate micro-/macro-architectures that once implemented, will improve the performance of a full-size implant and the rationale behind the design of the TMJ implant is outside the scope of this paper.

The general idea of the TMJ implant consists of a porous condylar head attached to the ramus via a fixture. A contact surface will facilitate bone marrow absorption to the porous condylar head and provides an environment for the native bone to regenerate as the implant degrades. The condylar head of the implant mimics the exact shape of the healthy joint and a hydrogel layer on top of the implant plays the cartilage role.

The full-size implant was ultimately designed with the surgery procedure in mind. That is, a major consideration was given to the method of implantation and how the design could be modified to facilitates the surgeon's job in the operating room. The surgery procedure in simple terms involves the opening of the skin to access the temporomandibular joint, resection of the damaged condylar head, and implantation of the device. Table 12 provides a graphical representation of the ramus, the cutting guide, and the TMJ implant in the surgery procedure sequence.

	Front	Isometric 1	Isometric 2
Full Ramus			
Resection		Cur	tting saw
Implantation			

Table 12. graphical view of the surgery procedure

Through practice surgeries, it was observed that two issues may arise with our initial design. The first was that the mandible might be resected at the wrong height or at the wrong angle. This will create a gap between the contact surface and the resected mandible and will halt the flow of the bone marrow to the condylar head and the hydrogel. The second issue was the thickness of the implant fixture. If too thin, the implant will bend at the intersection of the condylar head and the ramus fixture, and if too thick, it makes implantation more difficult and will stick out of the skin once implanted. These issues were resolved in the final design of the implant discussed in the next chapters.

The micro-architecture with the highest capillary forces and the micro-architecture with the strongest bone-to-cartilage bond were incorporated into the condylar head of the implant to expedite the rate of bone regeneration and to reinforce the hydrogel integration, respectively.

3.2 Materials and Methods

3.2.1 Design of the Implant

DICOM images from the patients' heads were acquired. These images were imported into Seg3D (CIBC) to create volumetric files representing the complete head. The condyle was resected with a sectioning plane that was referenced to four anatomical landmarks, two inferior orbits, and two superior external auditory meatuses, resulting in a consistently resected condylar head across all patients. These files were then imported into Meshmixer (Autodesk) to design the implant and the corresponding cutting guide.

The implant design consists of the condylar head and the ramus fixture. The condylar head is the exact geometry that was extracted from the DICOM images, and the ramus fixture is an extension of the ramus that attaches to the resected ramus. The cutting guide consists of the ramus fixture and an extended plate in place of the resected condylar head that is intended to help the surgeon with cutting the mandible at the right height and angle.

	Front	Back	Isometric 1	Isometric 2
Implan t				
Cuttin g Guide				

Table 13. graphical views of the implant and the cutting guide

A region of the ramus expanding from the condylar head to the angle of the mandible was selected to be extruded out in the inferior and posterior directions. This creates a pocket for the ramus to slide into and facilitates the alignment of the Implant and the ramus during the surgery. The extruded volume was tapered off towards the bottom of the implant so that it is less visible when implanted. Countersink pinholes were inserted normal to the surface of the implant and the cutting guide to affix them to the ramus. The location of these holes was selected such that they will be easily accessible to the surgeon through the incisions that are made on the side of the head. Right at the bottom of the condylar head, a plate wrapping around the mandible was added to the

cutting guide fixture to help the surgeon align the cutting tool parallel to the contact surface. This plate was lowered in height to account for the thickness of the cutting tool (table 13).

On the condylar head of the implant, several features were implemented to reinforce the bone-to-cartilage integration. A region covering the top surface of the condyle (approximating the region of contact between bone and cartilage) was sunken to create a seamless curve when the hydrogel is deposited on the implant. This reduces the chances of the hydrogel being sheared off when the jaw is being used. The deep slot design studied in <u>chapter 2.3.2</u> was implemented inside the sunken region. All of the features considered in the design of the implant are shown in figure 3.1.



Figure 3.1 Features of the TMJ implant.

3.2.2 Fabrication of the implant

Since the contact surface plays an integral role in the transportation of the bone marrow, its quality is of high importance and requires this surface to be printed without any support.

To determine the maximum overhang printable with our material, an overhang calibration sample was downloaded from <u>Thingiverse</u>. From the printed part it was observed that up to 50° overhang produces satisfactory results. The implant was then oriented on the print bed such that the contact surface had a lower overhang angle than 50° .

To minimize its use, support structures were manually created only for areas where the overhang was more than 50 degrees. This included the bottom of the implant and the posterior side of the condylar head.

To impose different micro-architectures in the implant, the condylar head shape was extracted from the implant model in Meshmixer (Autodesk) and was used as a mask to define new settings for the condyle internal structure. The print settings have been determined in the previous chapter based on the results of the capillary action and bone-to-cartilage studies. These parameters were used in the "Per Model" settings of Cura to create a porous micro-architecture where the condylar head mask meets the implant.

The final print configuration of the implant consists of three elements: the shell, the infill, and the support structures. The shell (colored red), as the name suggests, refers to the outer wall of the print models and could be seen in the ramus fixture and around the condylar head (figure 3.2, C). The infill (colored orange) refers to the structures that are located inside the shell (figure 3.2, A). The support structures (colored blue) are the pillars helping the implant to maintain its shape where the overhang is larger than 50 degrees (figure 3.2, B).



Figure 3.2 Implant print elements.

3.3 Results and Discussion

Transition to the full-size implant proved to accompany some difficulties along the way. One of the issues that arose when fabricating the implant was the instability of the print as it reached a higher height. The thin and tall ramus fixture made it proportionally more difficult to print the implant with the increase of height to the point where the print would fail. The addition of the support structure to the bottom of the implant and the thickening of the fixture were two modifications that helped stabilize the implant print.

Another technique used in the fabrication of the implant was postprocessing of the gcode to achieve different layer heights in a single print job. This is helpful in two ways; first, it was determined in <u>chapter 2.2.2</u> that our filament prints optimally at 10mm/s for the shell and the support structure and at 2.5mm/s for the infill. This means that printing a single implant could take up to 6 hours if the whole implant were to be printed with the layer height that proved to be most beneficial to the capillary forces in the condylar head. Changing the layer height to a higher value

in the ramus fixture could save up to 2.5 hours of print time. Second, we can change the mechanical properties of the implant by changing the layer height[80][81]. It also could be investigated whether different layer heights have a different rate of degradation. This means we can tailor our structure to accommodate the rate of degradation we need in certain locations.

In the initial implant design, the condylar head consisted of only the infill structure. This was later changed to the current design where a shell wraps around the porous infill structure to provide a smoother surface for the condylar head. This shell has only one perimeter and allows the porous structure to extend to the edges of the condylar head so as to not reduce its capacity for bone regeneration. The sunken region on top of the condylar head was expanded in all directions in order to cover as much area as possible. The extent of this expansion, however, was limited by the print orientation and the design of the implant.

4. Conclusion

4.1 Summary

In the first chapter of this paper, we talked about the current treatment options for temporomandibular disorders and mentioned instances where it falls short. We identified a gap in the clinical solution to the total joint replacement of the temporomandibular joint and pointed to several groups of people who would benefit from a novel approach. It was then discussed that tissue engineering and additive manufacturing have the potential to provide a new strategy to tackle this problem.

It was proposed that a biodegradable bi-phasic TMJ implant could replace the metal implants that are the current standard treatment options and overcome some of the limitations of metal implants. An implant design was conceptualized, and several features were proposed to improve the functionality of the implant.

This paper is divided into two phases; Micro-Architecture, where we design and fabricate structures on a micro level, and Macro-Architecture, where we utilized our findings from phase one to design an improved full-size implant.

In phase one, we started by preparing materials for our studies. This included a filament that was used to print the bone region of the implant and a hydrogel used to print the cartilage region. Through experiment, the optimal print settings for these materials were determined.

Several microscopic features were proposed and studied on standardized test samples to determine their effects. The first group of features was intended to increase the uptake capacity of the implant which in turn could result in a faster and higher rate of bone regeneration. The second group of features was intended to reinforce the bond between the bone and the cartilage regions and keep the hydrogel intact with the scaffolding of the implant.

In phase two, a full-size TMJ implant was designed. The micro-architectures that proved to be most beneficial to the performance of the test samples were incorporated in the macroarchitecture. Along with the implant, a cutting guide was designed that helps the surgeon to position the implant correctly during the surgery.

4.2 Contribution

In this paper, we proved that by modifying the design and print parameters of micro-/macroarchitectures we can tailor their responses to different factors and improve their performance. In <u>chapter 2.3.1</u> we showed that the test sample with 0.6 mm line distance and 0.12 mm layer height absorbed more solution compared to other designs, and in <u>chapter 2.3.2</u> we determined that the geometry with the deep slot had the strongest bond between bone and cartilage regions.

In <u>chapter 3.2.1</u> and <u>chapter 3.2.2</u>, we proposed a design and the fabrication process for a full-size implant and a cutting guide that could be a baseline for all future investigations on the design and performance of this implant.

4.3 Future Works

As mentioned before, the studies in <u>chapter 2.2.3</u> were performed with colored PBS to determine the capillary forces induced by several micro-architectures. It is possible that the results of those experiments would be different if the solution is changed, so it would be interesting to repeat those experiments with bone marrow to determine whether the results of the study would change.

It is also interesting to find the actual capacity of the micro-architectures proposed in <u>chapter</u> 2.2.3 by repeating the study at bigger scales. It was observed that the results of the capillary action studies were limited by the volume of the test samples and the size of their wells.

In the bone-to-cartilage integration studies, it was noticed that the hydrogels that failed, mostly did so in locations where there were sharp edges, for example, in the corners of slots and anchors. A future study could characterize the response of the hydrogel to axial force when these corners are less sharp and less tension is stored in them.

Another domain that could be further investigated is the orientation of the implant on the print bed, which will affect the mechanical properties, and possibly the degradation rate of the implant. This could also provide new opportunities for the design and incorporation of new micro-architectures which are impossible to implement in the current orientation. With the use of soluble materials, it is possible to find other orientations for the print in which the performance of the implant improves.

5. References

- [1] N. A. Vizniak *et al.*, "Muscle manual," p. 595, 2002.
- [2] L. Guarda-Nardini, D. Manfredini, and G. Ferronato, "Temporomandibular joint total replacement prosthesis: current knowledge and considerations for the future," *International Journal of Oral and Maxillofacial Surgery*, vol. 37, no. 2. Churchill Livingstone, pp. 103–110, Feb. 01, 2008, doi: 10.1016/j.ijom.2007.09.175.
- [3] Temporomandibular Joint Total Joint Replacement TMJ TJR. 2016.
- [4] J. J. Buescher, "Temporomandibular Joint Disorders," Nov. 2007. Accessed: Dec. 18, 2020.[Online]. Available: www.aafp.org/afp.
- [5] C. H. Wilkes, "Internal Derangements of the Temporomandibular Joint: Pathological Variations," Arch. Otolaryngol. Neck Surg., 1989, doi: 10.1001/archotol.1989.01860280067019.
- [6] R. de Leeuw, "Internal Derangements of the Temporomandibular Joint," *Oral and Maxillofacial Surgery Clinics of North America*. 2008, doi: 10.1016/j.coms.2007.12.004.
- [7] E. F. Wright and S. L. North, "Management and Treatment of Temporomandibular Disorders: A Clinical Perspective," *Journal of Manual and Manipulative Therapy*, vol. 17, no. 4. Maney Publishing, pp. 247–254, 2009, doi: 10.1179/106698109791352184.
- [8] R. Gray, "Manual of temporomandibular disorders," *Br. Dent. J.*, vol. 199, no. 6, pp. 399–399, Sep. 2005, doi: 10.1038/sj.bdj.4812802.
- [9] R. N. Arm, "Orofacial Pain. Guidelines for Assessment, Diagnosis, and Management," *Implant Dent.*, vol. 5, no. 4, p. 304, 1996, doi: 10.1097/00008505-199600540-00043.
- [10] B. Bagis, E. Aydogan Ayaz, S. Turgut, R. Durkan, and M. Özcan, "Gender Difference in Prevalence of Signs and Symptoms of Temporoman-dibular Joint Disorders: A Retrospective Study on 243 Consecutive Patients," *Int. J. Med. Sci*, vol. 9, no. 7, pp. 539– 544, 2012, doi: 10.7150/ijms.4474.
- [11] L. Leresche, "EPIDEMIOLOGY OF TEMPOROMANDIBULAR DISORDERS: IMPLICATIONS FOR THE INVESTIGATION OF ETIOLOGIC FACTORS."

- [12] A. Aryaei, N. Vapniarsky, J. C. Hu, and K. A. Athanasiou, "Recent Tissue Engineering Advances for the Treatment of Temporomandibular Joint Disorders," *Curr. Osteoporos. Rep.*, 2016, doi: 10.1007/s11914-016-0327-y.
- [13] U. Onoriobe, M. Miloro, C. Sukotjo, L. G. Mercuri, A. Lotesto, and R. Eke, "How Many Temporomandibular Joint Total Joint Alloplastic Implants Will Be Placed in the United States in 2030?," *J. Oral Maxillofac. Surg.*, vol. 74, no. 8, pp. 1531–1538, Aug. 2016, doi: 10.1016/j.joms.2016.04.011.
- [14] W. F. T. Lai, J. Bowley, and J. G. Burch, "Evaluation of Shear Stress of the Human Temporomandibular Joint Disc," J. Orofac. Pain, 1998.
- [15] M. Wieckiewicz, K. Boening, P. Wiland, Y. Y. Shiau, and A. Paradowska-Stolarz, "Reported concepts for the treatment modalities and pain management of temporomandibular disorders," *Journal of Headache and Pain*, vol. 16, no. 1. Springer-Verlag Italia s.r.l., pp. 1–12, Dec. 01, 2015, doi: 10.1186/s10194-015-0586-5.
- [16] M. Wieckiewicz, K. Boening, P. Wiland, Y.-Y. Shiau, and A. Paradowska-Stolarz, "Reported concepts for the treatment modalities and pain management of temporomandibular disorders," 2011, doi: 10.1186/s10194-015-0586-5.
- [17] R. A. Meyers, K. P. Schellhas, H. D. Hall, A. T. Indresano, and C. H. Wilkes, "Guidelines for diagnosis and management of disorders involving the temporomandibular joint and related musculoskeletal structures. American Society of Temporomandibular Joint Surgeons.," *Northwest Dent.*, vol. 71, no. 5, pp. 21–27, 1992.
- [18] P. A. Toller, "Osteoarthrosis of the mandibular condyle," Br. Dent. J., 1973, doi: 10.1038/sj.bdj.4802982.
- [19] E. Tanaka, M. S. Detamore, and L. G. Mercuri, "Degenerative disorders of the Temporomandibular joint: etiology, diagnosis, and treatment," *Journal of Dental Research*, vol. 87, no. 4. pp. 296–307, Apr. 2008, doi: 10.1177/154405910808700406.
- [20] S. G. G. Richard A. Pertes, "Overview of physical therapy modalities and procedures," in *Clinical Management of Temporomandibular Disorders and Orofacial Pain*, Quintessence Pub, 1995.

- [21] M. L. McNeely, S. A. Olivo, and D. J. Magee, "A systematic review of the effectiveness of physical therapy interventions for temporomandibular disorders," *Physical Therapy*. 2006, doi: 10.1093/ptj/86.5.710.
- [22] W. Shankland, "Temporomandibular disorders: standard treatment options.," *undefined*, 2004.
- [23] D. W. Nitzan, "Arthrocentesis-Incentives for Using This Minimally Invasive Approach for Temporomandibular Disorders," *Oral and Maxillofacial Surgery Clinics of North America*, vol. 18, no. 3. Elsevier, pp. 311–328, Aug. 01, 2006, doi: 10.1016/j.coms.2006.03.005.
- [24] P. A. Toller, "Use and Misuse of Intra-Articular Corticosteroids in Treatment of Temporomandibular Joint Pain," J. R. Soc. Med., 1977, doi: 10.1177/003591577707000704.
- [25] D. W. Nitzan and A. Price, "The use of arthrocentesis for the treatment of osteoarthritic temporomandibular joints," J. Oral Maxillofac. Surg., 2001, doi: 10.1053/joms.2001.26716.
- [26] D. Diraçoğlu *et al.*, "Arthrocentesis versus nonsurgical methods in the treatment of temporomandibular disc displacement without reduction," *Oral Surgery, Oral Med. Oral Pathol. Oral Radiol. Endodontology*, 2009, doi: 10.1016/j.tripleo.2009.01.005.
- [27] C. Linical and P. Ractice, "Internal Derangements of the Temporomandibular Joint: The Role of Arthroscopic Surgery and Arthrocentesis," J. Can. Dent. Assoc. (Tor)., 2000.
- [28] E. A. Al-Moraissi, "Arthroscopy versus arthrocentesis in the management of internal derangement of the temporomandibular joint: A systematic review and meta-analysis," *International Journal of Oral and Maxillofacial Surgery*, vol. 44, no. 1. Churchill Livingstone, pp. 104–112, Jan. 01, 2015, doi: 10.1016/j.ijom.2014.07.008.
- [29] M. G. Koslin, A. T. Indresano, and L. G. Mercuri, "Temporomandibular Joint Surgery," J. Oral Maxillofac. Surg., vol. 70, no. 11, Supplement 3, pp. e204–e231, 2012, doi: https://doi.org/10.1016/j.joms.2012.07.036.
- [30] J. D. Wagner and E. L. Mosby, "Assessement of Proplast-Teflon disc replacements," J. Oral Maxillofac. Surg., vol. 48, no. 11, pp. 1140–1144, Nov. 1990, doi: 10.1016/0278-

2391(90)90528-A.

- [31] N. De Meurechy and M. Y. Mommaerts, "Alloplastic temporomandibular joint replacement systems: a systematic review of their history," *International Journal of Oral and Maxillofacial Surgery*, vol. 47, no. 6. Churchill Livingstone, pp. 743–754, Jun. 01, 2018, doi: 10.1016/j.ijom.2018.01.014.
- [32] W. L. McCarty and W. B. Farrar, "Surgery for internal derangements of the temporomandibular joint," *J. Prosthet. Dent.*, 1979, doi: 10.1016/0022-3913(79)90174-4.
- [33] R. A. Williamson, D. McNamara, and W. McAuliffe, "True eminectomy for internal derangement of the temporomandibular joint," *Br. J. Oral Maxillofac. Surg.*, 2000, doi: 10.1054/bjom.2000.0467.
- [34] B. L. Florine, D. J. Gatto, M. L. Wade, and D. E. Waite, "Tomographic evaluation of temporomandibular joints following discoplasty or placement of polytetrafluoroethylene implants," J. Oral Maxillofac. Surg., 1988, doi: 10.1016/0278-2391(88)90080-8.
- [35] S. J. McKenna, "Discectomy for the treatment of internal derangements of the temporomandibular joint," *J. Oral Maxillofac. Surg.*, 2001, doi: 10.1053/joms.2001.26682.
- [36] M. K. Murphy, R. F. Macbarb, M. E. Wong, and K. A. Athanasiou, "Temporomandibular Joint Disorders: A Review of Etiology, Clinical Management, and Tissue Engineering Strategies HHS Public Access," 2013.
- [37] D. T. Brown and E. L. Gaudet, "Temporomandibular Disorder Treatment Outcomes: Second Report of a Large-Scale Prospective Clinical Study," *Cranio*, 2002, doi: 10.1080/08869634.2002.11746216.
- [38] G. A. Loupasis, K. Stamos, P. G. Katonis, G. Sapkas, D. S. Korres, and G. Hartofilakidis,
 "Seven- to 20-year outcome of lumbar discectomy," *Spine (Phila. Pa. 1976).*, 1999, doi: 10.1097/00007632-199911150-00005.
- [39] M. S. Detamore and K. A. Athanasiou, "Motivation, Characterization, and Strategy for Tissue Engineering the Temporomandibular Joint Disc," *Tissue Engineering*. 2003, doi: 10.1089/10763270360727991.

- [40] K. Ahtiainen *et al.*, "Autologous adipose stem cells and polylactide discs in the replacement of the rabbit temporomandibular joint disc," *J. R. Soc. Interface*, 2013, doi: 10.1098/rsif.2013.0287.
- [41] K. D. Allen and K. A. Athanasiou, "Scaffold and growth factor selection in temporomandibular joint disc engineering," J. Dent. Res., 2008, doi: 10.1177/154405910808700205.
- [42] M. S. Detamore and K. A. Athanasiou, "Evaluation of three growth factors for TMJ disc tissue engineering," Ann. Biomed. Eng., 2005, doi: 10.1007/s10439-005-1741-y.
- [43] C. Moura, D. Trindade, M. Vieira, L. Francisco, D. F. Angelo, and N. Alves, "Multi-Material Implants for Temporomandibular Joint Disc Repair: Tailored Additive Manufacturing Production," *Front. Bioeng. Biotechnol.*, 2020, doi: 10.3389/fbioe.2020.00342.
- [44] B. N. Brown *et al.*, "Inductive, Scaffold-Based, Regenerative Medicine Approach to Reconstruction of the Temporomandibular Joint Disk," *J. Oral Maxillofac. Surg.*, vol. 70, no. 11, pp. 2656–2668, Nov. 2012, doi: 10.1016/J.JOMS.2011.12.030.
- [45] K. Legemate, S. Tarafder, Y. Jun, and C. H. Lee, "Engineering Human TMJ Discs with Protein-Releasing 3D-Printed Scaffolds," *J. Dent. Res.*, vol. 95, no. 7, pp. 800–807, Apr. 2016, doi: 10.1177/0022034516642404.
- [46] J. R. Salash *et al.*, "Potential Indications for Tissue Engineering in Temporomandibular Joint Surgery," *Journal of Oral and Maxillofacial Surgery*, vol. 74, no. 4. W.B. Saunders, pp. 705–711, Apr. 01, 2016, doi: 10.1016/j.joms.2015.11.008.
- [47] D. E. Poswillo, "Biological reconstruction of the mandibular condyle," Br. J. Oral Maxillofac. Surg., 1987, doi: 10.1016/0266-4356(87)90003-9.
- [48] R. B. MacIntosh and F. A. Henny, "A spectrum of application of autogenous costochondral grafts," J. Maxillofac. Surg., 1977, doi: 10.1016/S0301-0503(77)80120-3.
- [49] R. B. MacIntosh, "The use of autogenous tissues for temporomandibular joint reconstruction," J. Oral Maxillofac. Surg., 2000, doi: 10.1016/S0278-2391(00)80019-1.

- [50] G. Obeid, S. A. Guttenberg, and P. W. Connole, "Costochondral grafting in condylar replacement and mandibular reconstruction," J. Oral Maxillofac. Surg., 1988, doi: 10.1016/0278-2391(88)90079-1.
- [51] N. Samman, L. K. Cheung, and H. Tideman, "Overgrowth of a costochondral graft in an adult male," *Int. J. Oral Maxillofac. Surg.*, 1995, doi: 10.1016/S0901-5027(05)80484-9.
- [52] T. Peltomäki, "Growth of a costochondral graft in the rat temporomandibular joint," J. Oral Maxillofac. Surg., 1992, doi: 10.1016/0278-2391(92)90278-8.
- [53] T. Peltomäki, K. Vähätalo, and O. Rönning, "The effect of a unilateral costochondral graft on the growth of the marmoset mandible," *J. Oral Maxillofac. Surg.*, 2002, doi: 10.1053/joms.2002.35729.
- [54] M. L. Pinson, C. A. Coop, and C. N. Webb, "Metal hypersensitivity in total joint arthroplasty," Annals of Allergy, Asthma and Immunology. 2014, doi: 10.1016/j.anai.2014.05.012.
- [55] L. Wang and M. S. Detamore, "Review Tissue Engineering the Mandibular Condyle," doi: 10.1089/ten.2006.0152.
- [56] T. M. Acri *et al.*, "Tissue Engineering for the Temporomandibular Joint," *Adv. Healthc. Mater.*, vol. 8, no. 2, p. 1801236, Jan. 2019, doi: 10.1002/adhm.201801236.
- [57] R. Schek, J. Taboas, S. Hollister, and P. Krebsbach, "Tissue engineering osteochondral implants for temporomandibular joint repair," *Orthod. Craniofac. Res.*, vol. 8, no. 4, pp. 313–319, Nov. 2005, doi: 10.1111/J.1601-6343.2005.00354.X.
- [58] K. Ueki *et al.*, "The use of polylactic acid/polyglycolic acid copolymer and gelatin sponge complex containing human recombinant bone morphogenetic protein-2 following condylectomy in rabbits," *J. Cranio-Maxillofacial Surg.*, vol. 31, no. 2, pp. 107–114, Apr. 2003, doi: 10.1016/S1010-5182(02)00187-7.
- [59] N. H. Dormer, K. Busaidy, C. J. Berkland, and M. S. Detamore, "Osteochondral interface regeneration of rabbit mandibular condyle with bioactive signal gradients," *J. Oral Maxillofac. Surg.*, vol. 69, no. 6, pp. e50–e57, Jun. 2011, doi: 10.1016/j.joms.2010.12.049.

- [60] J. P. Temple *et al.*, "Engineering anatomically shaped vascularized bone grafts with hASCs and 3D-printed PCL scaffolds," *J. Biomed. Mater. Res. - Part A*, 2014, doi: 10.1002/jbm.a.35107.
- [61] L. Ciocca *et al.*, "CAD-CAM-generated hydroxyapatite scaffold to replace the mandibular condyle in sheep: preliminary results.," *J. Biomater. Appl.*, vol. 28, no. 2, pp. 207–18, Aug. 2013, doi: 10.1177/0885328212443296.
- [62] D. Chen *et al.*, "Tissue engineered autologous cartilage-bone grafts for temporomandibular joint regeneration," *Sci. Transl. Med.*, vol. 12, no. 565, p. 6683, Oct. 2020, doi: 10.1126/SCITRANSLMED.ABB6683.
- [63] T. L. Aghaloo, P. K. Moy, and E. G. Freymiller, "Bone graft in the shape of human mandibular condyle reconstruction via seeding marrow-derived osteoblasts into porous coral in a nude mice model," *J. Oral Maxillofac. Surg.*, vol. 60, no. 10, pp. 1155–1159, Oct. 2002, doi: 10.1053/joms.2002.34991.
- [64] D. Howlader *et al.*, "Hydroxyapatite collagen scaffold with autologous bone marrow aspirate for mandibular condylar reconstruction," *J. Cranio-Maxillofacial Surg.*, vol. 45, no. 9, pp. 1566–1572, Sep. 2017, doi: 10.1016/j.jcms.2017.06.022.
- [65] Z. Wang, W. Xi, H. Zhu, and J. Yan, "Preparation and characterization of integrated condylar biomimetic scaffolds: A pilot study," *J. Hard Tissue Biol.*, 2016, doi: 10.2485/jhtb.25.89.
- [66] I. Gibson, D. W. Rosen, and B. Stucker, *Additive manufacturing technologies: Rapid prototyping to direct digital manufacturing*. 2010.
- [67] J. Giannatsis and V. Dedoussis, "Additive fabrication technologies applied to medicine and health care: A review," *Int. J. Adv. Manuf. Technol.*, 2009, doi: 10.1007/s00170-007-1308-1.
- [68] T. Baden, A. M. Chagas, G. Gage, T. Marzullo, L. L. Prieto-Godino, and T. Euler, "Open Labware: 3-D Printing Your Own Lab Equipment," *PLoS Biol.*, 2015, doi: 10.1371/journal.pbio.1002086.
- [69] S. Singare et al., "Rapid prototyping assisted surgery planning and custom implant design,"

Rapid Prototyp. J., 2009, doi: 10.1108/13552540910925027.

- [70] J. Y. Lee, B. Choi, B. Wu, and M. Lee, "Customized biomimetic scaffolds created by indirect three-dimensional printing for tissue engineering," *Biofabrication*, vol. 5, no. 4, p. 045003, Dec. 2013, doi: 10.1088/1758-5082/5/4/045003.
- [71] Hiroyuki Tetsuka and S. Ryon Shin, "Materials and technical innovations in 3D printing in biomedical applications," *J. Mater. Chem. B*, vol. 8, no. 15, pp. 2930–2950, Apr. 2020, doi: 10.1039/D0TB00034E.
- S. Ding, B. Zou, P. Wang, and H. Ding, "Effects of nozzle temperature and building orientation on mechanical properties and microstructure of PEEK and PEI printed by 3D-FDM," *Polym. Test.*, vol. 78, p. 105948, Sep. 2019, doi: 10.1016/J.POLYMERTESTING.2019.105948.
- [73] Ż. Górecka, J. Idaszek, D. Kołbuk, E. Choińska, A. Chlanda, and W. Święszkowski, "The effect of diameter of fibre on formation of hydrogen bonds and mechanical properties of 3D-printed PCL," *Mater. Sci. Eng. C*, vol. 114, p. 111072, Sep. 2020, doi: 10.1016/J.MSEC.2020.111072.
- [74] Feng Li, N. P. Macdonald, R. M. Guijt, and M. C. Breadmore, "Increasing the functionalities of 3D printed microchemical devices by single material, multimaterial, and print-pause-print 3D printing," *Lab Chip*, vol. 19, no. 1, pp. 35–49, Dec. 2018, doi: 10.1039/C8LC00826D.
- [75] D. Auhl, F. Rohnstock, O. Löschke, K. Schäfer, P. Wang, and M. H. Wagner, "3D-printing quality in relation to melt flow and fusion behavior of polymer materials," *AIP Conf. Proc.*, vol. 2107, no. 1, p. 030004, May 2019, doi: 10.1063/1.5109498.
- [76] A. Pandzic, D. Hodzic, and A. Milovanovic, "Effect of infill type and density on tensile properties of pla material for fdm process," 2019, doi: 10.2507/30th.daaam.proceedings.074.
- [77] T. Yao, J. Ye, Z. Deng, K. Zhang, Y. Ma, and H. Ouyang, "Tensile failure strength and separation angle of FDM 3D printing PLA material: Experimental and theoretical analyses," *Compos. Part B Eng.*, vol. 188, p. 107894, May 2020, doi:

10.1016/J.COMPOSITESB.2020.107894.

- [78] S. J. Polak, "Microstructure-induced capillary force effects on penetration and release in hydroxyapatite bone scaffolds," 2012, Accessed: Jul. 15, 2021. [Online]. Available: https://www.ideals.illinois.edu/handle/2142/42348.
- [79] Z. Jiao, B. Luo, S. Xiang, H. Ma, Y. Yu, and W. Yang, "3D printing of HA / PCL composite tissue engineering scaffolds," *Adv. Ind. Eng. Polym. Res.*, vol. 2, no. 4, pp. 196–202, Oct. 2019, doi: 10.1016/J.AIEPR.2019.09.003.
- [80] S. R. Rajpurohit and H. K. Dave, "Analysis of tensile strength of a fused filament fabricated PLA part using an open-source 3D printer," *Int. J. Adv. Manuf. Technol. 2018 1015*, vol. 101, no. 5, pp. 1525–1536, Nov. 2018, doi: 10.1007/S00170-018-3047-X.
- [81] A. M. Sousa, A. C. Pinho, and A. P. Piedade, "Mechanical properties of 3D printed mouthguards: Influence of layer height and device thickness," *Mater. Des.*, vol. 203, p. 109624, May 2021, doi: 10.1016/J.MATDES.2021.109624.

6. Appendix

		Descrip	tive Statistics		
	D , 1	Dependent	Variable: Upta	ake	
LayerHeight	Etch	LineDistance	Mean	Std. Deviation	N
.12	no	.6	61.03333333	12.61995773	3
			3333300	8967747	
		.8	67.59999999	10.14297786	3
			9999950	6484772	
		1.0	62.49999999	8.373768566	3
			9999964	183314	
		Total	63.71111111	9.590417671	9
			1111070	358773	
	yes	.6	445.3333333	21.19654059	3
			33333300	8251676	
		.8	208.6666666	65.59186941	3
			66666700	4839300	
		1.0	215.8999999	154.9546062	3
			99999980	56154880	
		Total	289.9666666	144.1475459	9
			66666700	38180970	
	Total	.6	253.1833333	211.0672159	6
			33333340	92125740	
		.8	138.1333333	87.93185240	6
			33333330	0973970	
		1.0	139.1999999	129.1970897	6
			99999960	50504830	
		Total	176.8388888	152.8794263	18
			88888870	03633520	
.18	no	.6	41.13333333	4.940985056	3
			3333360	983492	
		.8	36.19999999	10.63343782	3
			9999970	6027862	
		1.0	25.03333333	5.085600587	3
			3333303	279099	
		Total	34.12222222	9.584724536	9
			2222210	701357	
	yes	.6	75.16666666	11.38083183	3
			6666740	8373293	
		.8	76.93333333	31.11468035	3
			3333340	0814112	

		1.0	74.53333333	5.704676444	3
			3333350	228316	
		Total	75.5444444	16.84362721	9
			4444470	5590420	
	Total	.6	58.15000000	20.22510815	6
			0000055	7930833	
		.8	56.56666666	30.49981420	6
			6666660	7084437	
		1.0	49.78333333	27.53974703	6
			3333330	3454535	
		Total	54.83333333	25.11819120	18
			3333360	5016455	
.30	no	.6	34.56666666	6.469415223	3
			6666606	444364	
		.8	79.39999999	18.92405876	3
			9999930	1270004	
		1.0	78.93333333	34.60236022	3
			3333280	7784100	
		Total	64.29999999	29.94419810	9
			9999940	2470544	
	yes	.6	204.9333333	147.7176134	3
			33333400	83745850	
		.8	131.2999999	54.50055045	3
			99999980	5935745	
		1.0	167.8000000	35.26003403	3
			00000040	2882050	
		Total	168.0111111	86.74758562	9
			11111180	1221250	
	Total	.6	119.7500000	132.1074524	6
			0000000	77140780	
		.8	105.3499999	46.25424304	6
			99999970	8611230	
		1.0	123.3666666	57.83962886	6
			66666650	0035670	
		Total	116.1555555	82.52499245	18
			55555540	1286960	
Total	no	.6	45.57777777	14.10086679	9
			///7754	/62/884	-
		.8	61.06666666	22.74747238	9
		4.0	6666630	7058730	-
		1.0	55.48888888	29.92830785	9
	_		8888845	5792830	
		Total	54.0444444	23.20777942	27
			4444410	0724970	

yes	.6	241.8111111	179.0438245	9
		11111160	54523780	
	.8	138.9666666	73.12357690	9
		66666700	9229500	
	1.0	152.7444444	100.9766694	9
		4444440	72595380	
	Total	177.8407407	129.6295498	27
		40740730	69735300	
Total	.6	143.6944444	159.2866541	18
		4444460	88649400	
	.8	100.0166666	66.07655319	18
		66666650	6674420	
	1.0	104.1166666	87.88355430	18
		66666650	7088990	
	Total	115.9425925	111.4058966	54
		92592600	66287670	

	Univariate Tests							
	Dependent Variable: Uptake							
		Sum of						
H	Etch	Squares	df	Mean Square	F	Sig.		
no	Contrast	1107.742	2	553.871	.172	.843		
	Error	116039.373	36	3223.316				
yes	Contrast	56099.034	2	28049.517	8.702	.001		
	Error	116039.373	36	3223.316				

Each F tests the simple effects of LineDistance within each level combination of the other effects shown. These tests are based on the linearly independent pairwise comparisons among the estimated marginal means.

Pairwise Comparisons Dependent Variable: Uptake

						97.5% Co	onfidence
			Mean			Interval for	Difference ^b
	(I)	(J)	Difference	Std.		Lower	Upper
Etch	LineDistance	LineDistance	(I-J)	Error	Sig. ^b	Bound	Bound
no	.6	.8	-15.489	26.764	.566	-78.091	47.113
		1.0	-9.911	26.764	.713	-72.513	52.691

	.8	.6	15.489	26.764	.566	-47.113	78.091
		1.0	5.578	26.764	.836	-57.024	68.180
	1.0	.6	9.911	26.764	.713	-52.691	72.513
		.8	-5.578	26.764	.836	-68.180	57.024
yes	.6	.8	102.844*	26.764	.000	40.243	165.446
		1.0	89.067*	26.764	.002	26.465	151.668
	.8	.6	-102.844*	26.764	.000	-165.446	-40.243
		1.0	-13.778	26.764	.610	-76.380	48.824
	1.0	.6	-89.067*	26.764	.002	-151.668	-26.465
		.8	13.778	26.764	.610	-48.824	76.380

Based on estimated marginal means

*. The mean difference is significant at the .025 level.

b. Adjustment for multiple comparisons: Least Significant Difference (equivalent to no adjustments).