

Orthocell

Advancing tissue repair and regeneration

Investor update

April 2018

Paul Anderson Managing Director



Corporate overview

Orthocell is a regenerative medicine company delivering breakthrough products that restore mobility and function

Share price performance



1. Excludes 12.1m unquoted warrants with exercise price \$0.58, expiry 19-Nov-2020 and 3.3m unquoted options with exercise prices ranging from \$0.51-\$0.65 and expiry dates between Feb-2019 and Jun-2020
2. Cash figure does not include cash from share purchase plan announced Dec-2017

Trading information

Share price (11-Apr-18)	A\$0.305
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Shares on issue ¹	110.0m
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Market capitalisation	A\$33.5m
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Cash (as at 31-Dec-17)	A\$4.9m ²
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Debt (as at 31-Dec-17)	-
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Enterprise value	A\$28.6m
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Top shareholders (as at Feb-18)

Stone Ridge Ventures – Associated with non-executive director	9.3%
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Ming Hao Zheng – CSO and founder	6.7%
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Paul Anderson – Managing director	6.4%
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Qi Xiao Zhou – Non-executive director	5.8%
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Australian Super – Superannuation and pension fund	4.2%
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Founders story

Orthocell's founders have extensive commercial and scientific experience in regenerative medicine, having previously demonstrated success in developing, commercialising and monetising cell therapy products



Paul Anderson

Managing Director

- **20+ years' experience** within the medical device and cellular therapy fields, with intimate knowledge of the regenerative medicine space
- **Former Managing Director and Executive Director of Verigen Australia** (prior to its sale to Genzyme in 2005)
- Extensive and proven experience in **commercialising medical technology, corporate transactions, establishing certified manufacturing facilities, and navigating regulatory pathways** for cell therapies



Professor Ming Hao Zheng¹

Chief Scientific Officer

Significant educational, scientific and professional experience within China¹

- **Work (current):** Chung Kong Lecturing Professor, First Affiliated Hospital, Zhejiang University
- **Previous experience:** Histopathologist, Lecturer & Tutor at Sun Yat-Sen University of Medical Sciences²; Consultant Histopathologist at The Huizhou People's Hospital²
- **Education:** Masters of Medicine in Pathology, Sun Yat-Sen University of Medical Sciences; Bachelor of Medicine, Shantou University
- **Advisor, Assessor, Reviewer:** Cheung-Kon Scholar Foundation, Ministry of Education; Chinese National Science Foundation
- **Honours/Awards:** Chang Jiang Scholar at Zhejiang University, Honorary Professor at Sun Yat-Sen University of Medical Sciences² and Shantou University

1. Experience only reflects selected China-related experience

2. Department of Pathology

Board of directors



Dr Stewart James Washer
Chairman

- 20+ years' CEO and board experience
- Commercialisation, capital markets and corporate advisory

Commercialisation
Corporate
Capital markets



Paul Anderson
Managing Director

- 20+ years' regenerative medicine experience
- Former MD of Verigen
- Extensive experience in commercialising emerging technologies

Regenerative medicine
Commercialisation
Regulatory, clinical
manufacturing



Matthew Callahan
Non-Executive Director

- Developed 3 FDA approved products
- Previous investment director of 2 VC firms (life science focus)
- Extensive corporate and IP experience

Corporate
Regulatory and clinical
IP and legal



Professor Lars Lidgren
Board Member

- World leading innovator in the orthopaedic space
- Entrepreneur and founder of multiple biotech companies (Scandimed, Bone Support, AMeC and GWS)

Regenerative medicine
Research & development



Mr Qi Xiao Zhou
Board Member

- 15+ years' in China as a senior business manager and executive
- Experience within public markets of Hong Kong, China and Taiwan

Corporate
Commercial



What is regenerative medicine?

*Relatively new field of medicine that seeks to **repair injured or diseased tissue** by harnessing the **body's own regenerative capabilities***

Replacing**, engineering or regenerating, **human cells**, tissues or organs **to restore or establish normal function



Why is regenerative medicine a promising field?

***Ageing population** and rising musculoskeletal disorders*

Demand for safe, efficient and cost effective treatments

*International regulatory bodies (e.g. FDA), **accelerating development** and access to safe and effective regenerative medicine therapies*



What is Orthocell's position in this space?

*Orthocell is a **world leading** regenerative medicine company with **novel, first in class, most advanced** portfolio of products*

Innovative products

Orthocell is a regenerative medicine company delivering breakthrough products that restore mobility and function



CelGro®

*Soft tissue reconstruction platform
Medical device*

- **CE Mark** - approved in Europe for bone and tissue
- **Superior** - clinical performance
- **Commercialisation** - key European markets
- **Multiple indications** - superior collagen medical device
- **Optimised** - manufacturing capabilities
- **Lack of innovation** - static market
- **Global partnering**



Ortho-ATI®

*Regeneration of tendons
Cell therapy*

- **First in class** - cell therapy for tendon repair
- **Significant unmet clinical need**
- **Global partnering** - major US collaboration partner
- **De-risked** - proven safe and effective (>500 patients treated)
- **TGA** licence to manufacture and treat patients in Australia, Singapore and Hong Kong
- **US regulatory focus** - process underway

Significant market opportunity

At the forefront of a large and growing market opportunity in regenerative medicine in the musculoskeletal space

Addressable markets^{1,2}



CelGro®

>US\$2.6bn



Ortho-ATI®

>US\$7.7bn

Total addressable market is estimated to be in excess of
US\$10 billion p.a.

1. Addressable markets include US, Japanese, European and Australian markets

2. Ortho-ATI® addressable market includes the following indications: tennis elbow, rotator cuff, gluteal, patellar, hamstring and achilles. CelGro® addressable market includes the following indications: dental, rotator cuff and nerve.

CelGro® - initial focus in bone regeneration

CelGro® is approved in the EU (CE Mark) for dental bone and soft tissue repair

CelGro® - a true regenerative medicine scaffold

- ✓ **Superior tissue repair** – unique regenerative medicine qualities
- ✓ **Superior handling** characteristics over existing products
- ✓ **Proprietary SMRT™** manufacturing process
- ✓ **CE Mark** – dental bone and soft tissue repair approved for use in the EU

Dr Brent Allan, Chief Investigator, dental bone regeneration trial:
“CelGro® is an exciting new product with clear advantages over available alternatives.”

Dental bone procedure - approved for use in the EU



1. Defect Site



2. Bone Graft



3. Apply CelGro®



4. Insert Crown

CelGro® - bone regeneration market opportunity

Ideally positioned to gain market traction in the rapidly growing and significant market of oral medicine

Very favourable market dynamics...

- **No product innovation** to the dental market globally
- Existing products have **inferior functionality and handling characteristics**
- **Strong demand** from dentists / surgeons
- Market leader generates €50m p.a. in EU alone



***CelGro® has superior clinical performance
and a significant market opportunity***



Significant addressable market¹

>US\$0.6bn p.a.

~1.5m procedures per year

1. US, Japanese, European and Australian markets

CelGro® - bone regeneration commercial pathway

Key development and marketing approval milestones for CelGro® achieved, the focus is now on EU market entry in bone and soft tissue repair

Achieved significant development milestones



Optimised manufacturing

- ✓ Quality and proprietary SMRT™ manufacturing processes
- ✓ Scalable infrastructure in place



Regulatory approval

- ✓ Clinically validated products, completed studies
- ✓ CelGro® approved for marketing and distribution in the EU (CE Mark)



Key opinion leaders in place

- ✓ Dr Simion
- ✓ Dr Luongo
- ✓ World leaders in oral medicine and targeted by competitors

Market entry – next steps



EU Market

- Supplementary marketing clinical data
- Secure distribution partnerships
- Initial sales
- KOL led adoption in key markets

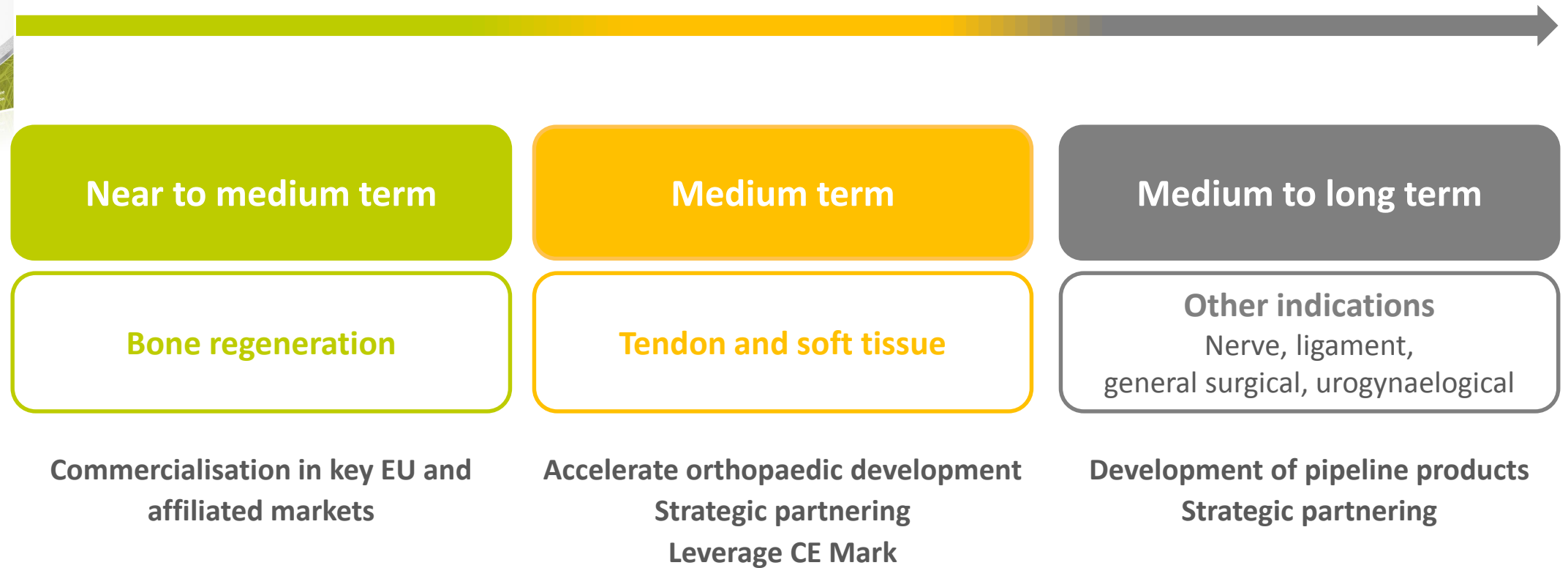


Expansion

- Leverage CE Mark for US and AU market entry
- Secure distribution partnerships
- M&A

CelGro® - strategic focus

Orthocell is driving market entry for bone repair, leveraging EU approval to accelerate introduction of the tendon and soft tissue indications



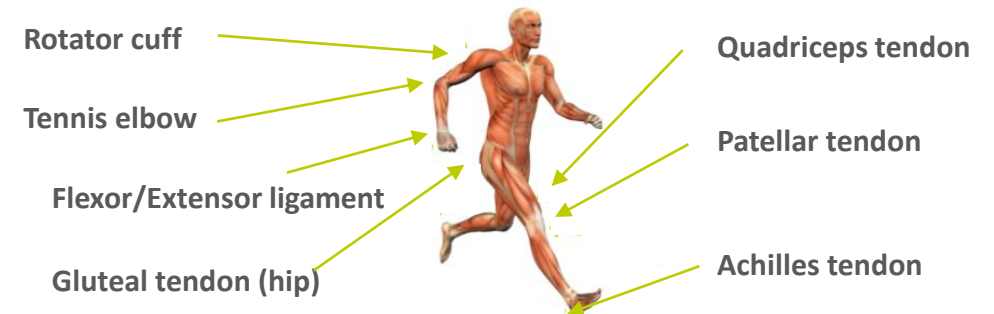
Chronic tendon injury – significant unmet clinical need

Chronic tendon injury is a common and painful disorder affecting millions of people every year with no effective, non-surgical treatments currently available

Significant unmet clinical need...

- **Millions of people suffer** from chronic tendon injury every year
- **Chronic tendon injury significantly reduces** the ability of patients to work, exercise, and perform routine daily activities
- **Increasing financial burden** on the public health care system expected as the population ages
- **Demand for new treatments** that are safe, minimally invasive (non surgical), effective and cost efficient

Multiple tendon injury sites...



There are no 'non-surgical' treatments currently available to treat chronic tendon injury

Ortho-ATI[®] - preferred therapy for tendon repair

Orthocell is currently focused on completing current clinical studies and preparing for US market entry

Ortho-ATI[®] overview

- **Breakthrough** in regenerative medicine
- Novel, cell therapy to **treat chronic degenerative tendon injuries**
- **Replenishes degenerative tissue** with healthy mature tendon cells, accelerating regeneration of tendon tissue
- Allows patients to **return to the, workplace, recreational activities and competitive sport**
- Extensive clinical validation - **over 500 patient implants** with Ortho-ATI[®] to date

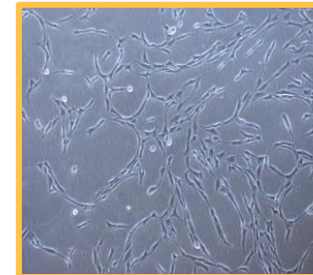
Two stage, minimally invasive procedure

1. Biopsy procedure



Healthy tendon cells removed via minimally invasive procedure

Tenocyte (cell) cultivation



Healthy cells grown at Orthocell's laboratory

2. Tenocyte (cell) implantation



Ultrasound guided implementation of healthy cells

4-5 week
end-to-end
process



Ortho-ATI® - multiple, large addressable markets

Ortho-ATI® is at the forefront of a large and growing market opportunity where the addressable market is estimated to be >US\$7.7bn p.a.

Estimated addressable market in excess of US\$7.7bn incorporates:

Tennis elbow¹

>US\$4.3bn

~1,000,000 procedures p.a.



Rotator cuff¹

>US\$2.4bn

~530,000 procedures p.a.



Other indications

>US\$1.0bn

Gluteal, Patellar, Hamstring & Achilles



Illustrative analysis: tennis elbow addressable market²

~864m

~11.3m p.a.

~1.2m p.a.

~1.0m p.a.

>US\$4.3bn

Total population

In key markets¹

Estimated incidence

Based on ~1.3%
incidence rate p.a.

Resistant patients

Assumed ~10.5% resistant to
conservative treatment

Total procedures

Assumed ~81.3% treatable
with Ortho-ATI®

Estimated market size

Assumed
US\$4,500 price

1. Includes US, EU, Japan, Australia and New Zealand markets

2. Illustrative analysis subject to rounding estimates

Ortho-ATI® - clinically validated product

Patients achieved a 207% improvement in grip strength following their treatment with Ortho-ATI®

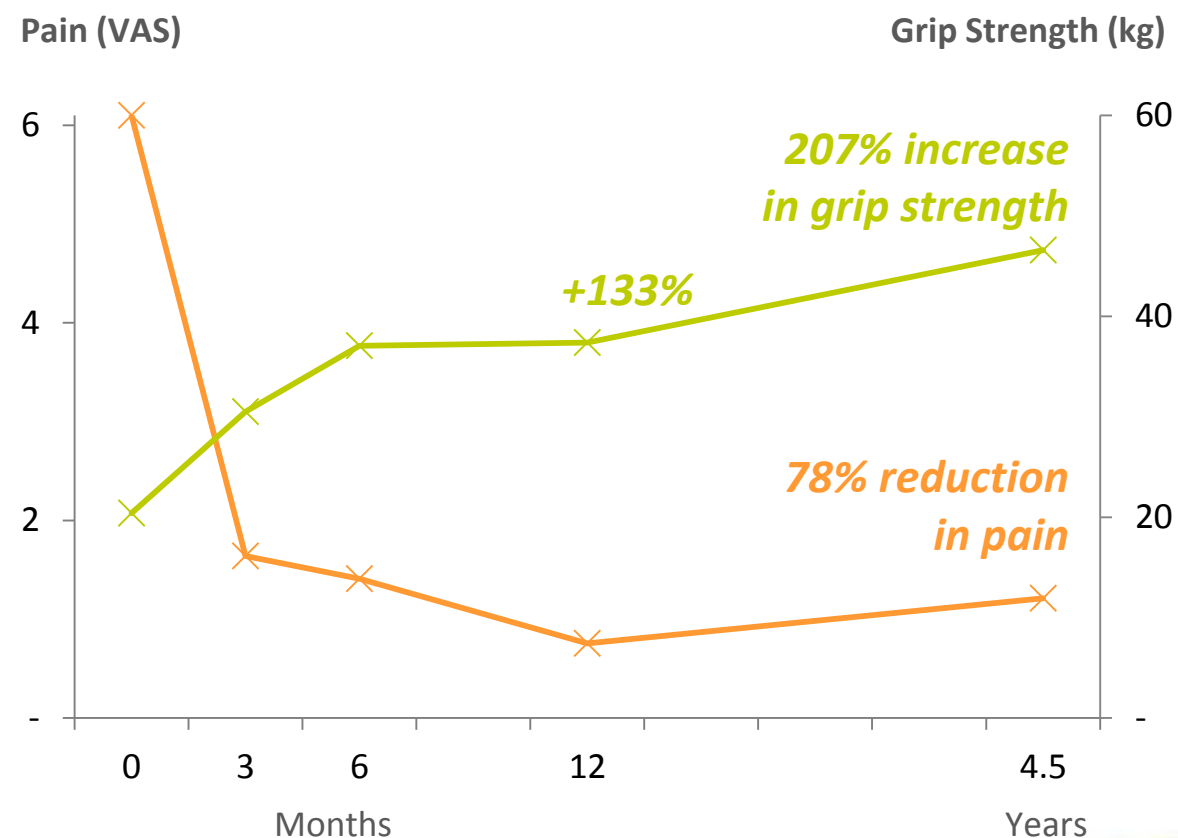
Ortho-ATI® tennis elbow study details

- Patients that had **failed all other treatments**
- The patients had been **experiencing symptoms for an average of 31 months**
- Data to 4.5 years published in *American Journal Sports Medicine*

207% increase in grip strength

Results from the trial are very encouraging; with clinical data showing patients are still pain-free four-and-a-half years after treatment.

Tennis elbow study results



Ortho-ATI® - patient testimonies

“I’m 100% happy with the Ortho-ATI® treatment, the only thing that enabled me to get back to doing normal daily activities without pain and return to swimming at an international level.”

Christian Sprenger
Swimming Olympic medallist



“I couldn’t write, drive, play my violin, or even walk without pain. Ortho-ATI® gave me my life back.”

Jasmine Trebse
Professional Violinist

Ortho-ATI® - research collaboration

Ortho-ATI® has attracted the interest of the worlds largest health care company

Key factors in attracting research collaboration interest

- ✓ **Large unmet clinical need**
1.5m+ addressable procedures per year in the shoulder and elbow alone
- ✓ **Clinical validation**
Published clinical data in American Journal of Sports Medicine
Completed 500+ patient implants to date
- ✓ **Optimised manufacturing capability**
GMP-certified and TGA-licensed facility¹
PPI release criteria in place²



Ortho-ATI® research collaboration

The objective of this study is to assess the effectiveness of Autologous Tenocyte Injection (Ortho-ATI®) compared to corticosteroid injection in the treatment of **rotator cuff tendinopathy** and tear. The trial is being undertaken in **collaboration with DePuy Synthes Products, Inc., part of the Johnson & Johnson Medical Device Companies**

1. GMP: good manufacturing practices; TGA: Therapeutic Goods Administration
2. PPI: purity, potency and identity

Significant upside value potential

Significant value upside potential exists given Orthocell's valuable diversified product portfolio of cell therapies and regenerative medicine products, when compared to similar global peers

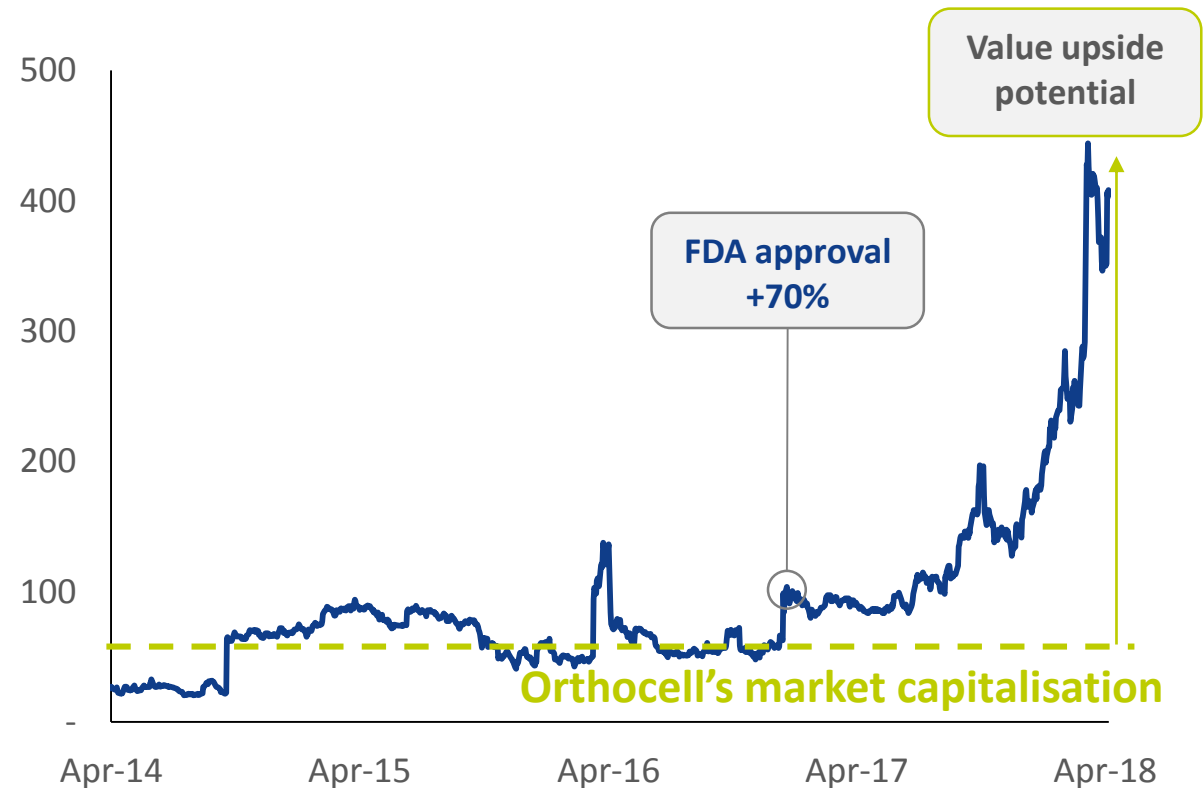


Develops, manufactures, and markets two regenerative medicine products for epidermal autografts and cartilage regeneration

Product comparison (Orthocell vs. Vericel)

	Orthocell	Vericel
ACI stem-cell product (cartilage focus)	✓	✓
ATI stem-cell product (tendon focus)	✓	✗
Platform technology (multi-use scaffold)	✓	✗

Vericel's market capitalisation (US\$m)



Significant M&A activity in regenerative medicine

Smith & Nephew's recent US\$210m acquisition of Rotation Medical, a comparable peer to Orthocell, demonstrates the strong interest by major players within the regenerative medicine sector

US\$210m acquisition of Rotation Medical



- **M&A activity** - major players are actively securing emerging regenerative medicine technologies
- **US\$210m acquisition** of Rotation Medical in October 2017
- **Rotation Medical** - medical scaffold company focused on rotator cuff repair with FDA 510(k) clearance and initial sales

Product comparison (Orthocell vs. Rotation Medical)

Key comparisons	Orthocell	Rotation Medical
Tendon regeneration	✓	✓
Platform technology	✓	✗
Multiple regenerative medicine products	✓	✗
Regulatory approval	✓	✓

Upcoming catalysts

Summary of the upcoming operational milestones



CelGro®

- Supplementary marketing clinical data release – April 2018
- Secure distribution networks - 2Q 2018
- First sales in the EU – 2Q 2018
- US and Australian market authorisation – 1Q 2019
- Strategic partnership discussions - ongoing

Ortho-ATI®

- Accelerate completion of rotator cuff study – 2Q 2018
- Pre-Investigation New Drug meeting with FDA – 3Q 2018
- Strategic partnership discussions - ongoing

Key investment highlights



Significant upside

*Significant market interest
Addressable markets worth
>US\$10bn p.a.*



De-risked product portfolio

*Substantial clinical data
CE Mark (EU) achieved for
CelGro¹*



Validated manufacturing process

*GMP-certified and TGA-
licensed manufacturing
capabilities*



Credentialed and highly aligned leadership team

*Proven track record in
commercialising cell therapy
products*

1. CelGro® has CE Mark (EU) for dental bone and soft tissue reconstructive applications

Appendix

Executive management, key product portfolio, intellectual property, additional product information, clinical data and R&D pipeline



Executive Management Team

Professor M.H. Zheng Chief Scientific Officer

- Inventor of the Orthocell technology with a strong track record of innovation
- Director of Research in the Department of Orthopaedic Surgery at the University of WA
- Research focus on new treatment methods for osteoporosis, osteoarthritis and tendon injuries using cellular and molecular biology techniques
- Ph.D., Doctor of Medicine and Fellow of the Royal College of Pathologists

Alexander McHenry Chief Operating Officer

- Over 14 years of experience in corporate advisory and management consulting
- Background in the implementation of corporate transaction and business transformation initiatives
- Master of Business Administration from The University of Western Australia
- Former senior manager of the strategy and operations division of Deloitte, Perth WA

Nicole Telford Chief Financial Officer

- Chartered accountant with over 14 years' commercial experience in financial controller and group accountant roles
- Background of broad commercial experience in financial and management reporting, office administration and staff management
- Achieved professional qualifications whilst employed with Arthur Andersen in the audit division

Gregor Maier Sales & Marketing Director

- Over 14 years' commercial experience in the pharmaceutical and surgical industries having previously worked in the UK, EU, Malaysia and South Africa
- Oversaw successful launch of Ortho-ATI® and Ortho-AC® launch into Hong Kong, NZ and Singapore markets
- Wealth of senior management experience following key management roles in Malaysia and South Africa

Dr Clair Lee Clinical Research Manager

- Clinical research professional with over 17 years of experience in pharmaceutical and medical device clinical trials
- Specialist in translation of basic scientific research into clinical trial and regulatory application strategies
- Ph.D. in Cell Biology from the University of WA
- Former Program Manager and Faculty Member at Telethon Kids Institute, Perth WA

Monique Cannon Quality Director

- Background includes research in cellular development and differentiation, with significant experience in cell culture
- Previous role in the GMP-compliant manufacture of an autologous cell-based therapeutic product
- Over 14 years experience in coordination of qualification systems, quality system management, and significant regulatory experience (ISO, TGA, FDA)

Key product portfolio

Diversified product portfolio with exciting R&D development pipeline

Diversified health care	Products	Key focus	Applications	Stages of development		
				Discovery	Pre-clinical	Clinical
Pure collagen medical device	CelGro®	Biological medical device for soft tissue reconstruction & repair	Bone	EU (CE Mark)		
			Orthopaedic			
			ACL ligament			
Cell based therapies	Ortho-ATI®	Tendon regeneration	Tennis elbow, gluteal etc.			
	Ortho-ACI®	Cartilage regeneration	Knee & ankle cartilage repair	AUS (ARTG)		
	R&D pipeline	"Off-the shelf" tissue repair therapies	Growth factors			
		Allogenic tendon repair	Lab grown tendon			

CelGro®

Tissue and bone repair

Off the shelf collagen scaffold to support soft tissue and bone repair



Ortho-ATI®

Regeneration of tendons

Two stage, minimally invasive, walk-in & walk-out procedure



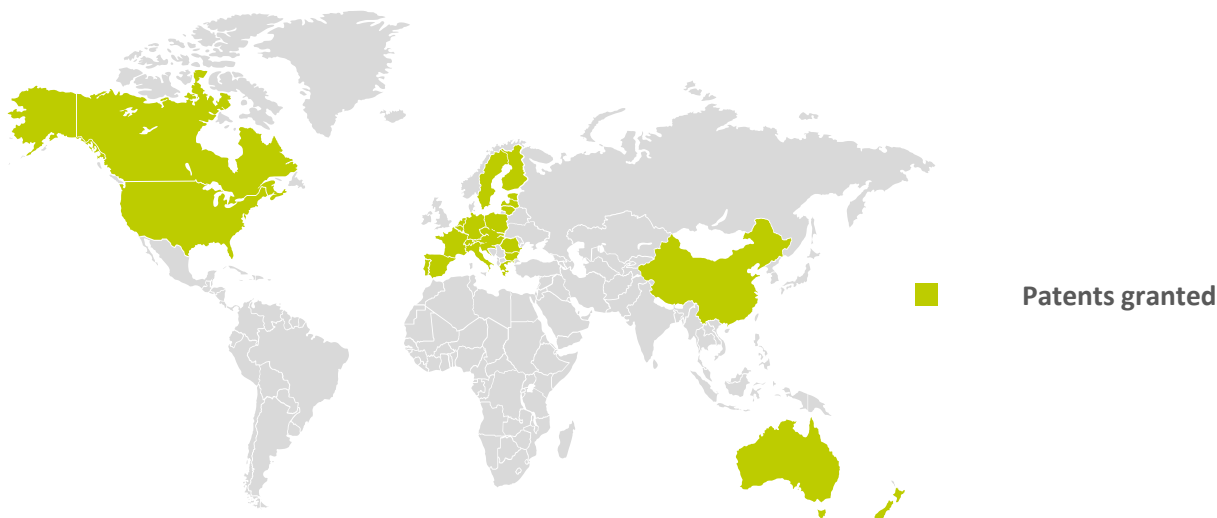
Proven and logical clinical development strategy, grounded on an evidence-based translational programme to deliver cost effective treatments that transform patients lives

Key products are well protected and clinically validated

Orthocell has significant levels of clinical validation and has protected its suite of development products through various patents granted across major jurisdictions, with further applications awaiting grant

IP portfolio and strategy

- Focused on maintaining patent protection for its leading manufacturing technologies and treatment processes
- Maintains an active program of patenting, with ownership of 27 granted patents across the Ortho-ATI®, Ortho-ACI® and CelGro® technologies, and related methods of treatment and manufacturing
- Currently has 31 patent applications across 7 different patent families
- Patent protection is targeted at major jurisdictions across Australia, Asia, Europe and North America, which represent key regions where Orthocell is targeting regulatory approval



Clinical studies

1,000+ Patients treated to dated

7 Results published in academic journal

CelGro® - key advantages

CelGro® boasts superior tissue repair qualities and can be customised to accommodate the specific needs of tissue repair in multiple applications, making it the best in class bioderived scaffold on the market



Orthocell's **proprietary SMRT™ manufacturing process** method produces collagen scaffolds with **numerous competitive advantages** over existing tissue repair scaffolds, particularly in the areas of **compatibility, tensile strength, promotion of quality tissue repair and versatility**

Key advantages	Dental		Tendon		Nerve			Comments
	Biogide®	BioMed Extend®	GraftJacket®	Tissuemend®	AxoGuard®	Neuroflex®		
Proven compatibility	✓	+/-	+/-	+/-	+/-	+/-	✓	SMRT™ process removes all DNA and other tissue components, collagen scaffold is biocompatible and naturally degrades after implantation at the same rate as the body heals
Strong mechanical properties	+/-	✓	✓	✓	✓	✓	✓	No cross-linking or artificial additives required to boost strength of intact collagen fibres. CelGro® is easy to use, ductile and very strong
Quality of tissue repair (Native bilayer structure)	✓	x	x	x	x	x	✓	Preservation of the natural collagen structure from the source material promotes healing and regeneration. CelGro® integrates with the soft tissue under repair leaving no remnant material to scar or cause inflammation
Versatile/customisable to multiple applications	x	x	x	x	x	x	✓	Easily customisable, with multiple applications and the ability to be used on its own in surgical repair, or in combination with stem cells and other drugs

Biogide® currently generates €50m p.a. in dental bone graft sales in the EU alone – CelGro® has been proven, through clinical studies, to produce superior bone regeneration

+/- No clinical data available to support characteristic required for optimal tissue repair

CelGro® - platform technology

CelGro® is a platform technology that supports tissue growth and enables rapid repair, it has demonstrated clinical efficacy and has a clear path to market across multiple applications

Multiple applications

Bone

e.g. Dental surgery



1. Defect Site

Insufficient bone volume to anchor the dental implant



2. Bone Graft

Defect site filled with bone grafting material



3. Apply CelGro®

Customised to size and placed over defect site to contain graft, 'guiding' bone regeneration



4. Implant Crown

Gum tissue stitched over CelGro® and Crown secured

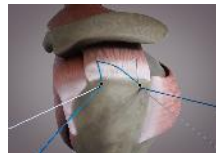
Tendon

e.g. Orthopaedic surgery



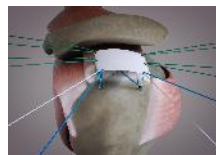
1. Tendon Tear

Tendon detached from bone and retracted at tear site



2. Preparation of Repair Site

Torn tendon trimmed and sutures anchored in healthy bone



3. Apply CelGro®

Customised to size and placed over repair site to promote tendon healing



4. Reattachment

Tendon and CelGro® secured in place with sutures

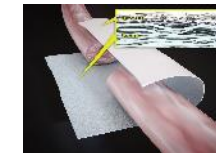
Neurological

e.g. Peripheral nerve repair



1. Peripheral Nerve Injury

Nerve severed and ends retracted into wound after traumatic injury to limb



2. Preparation of Repair Site

Nerve ends located and brought to repair site without tension



3. Microsurgical Repair with CelGro®

CelGro® secured around nerve ends, forming a sealed conduit



4. Nerve Healing

Conduit protects nerve ends and enhances regeneration

CelGro® - clinical trials completed and in progress

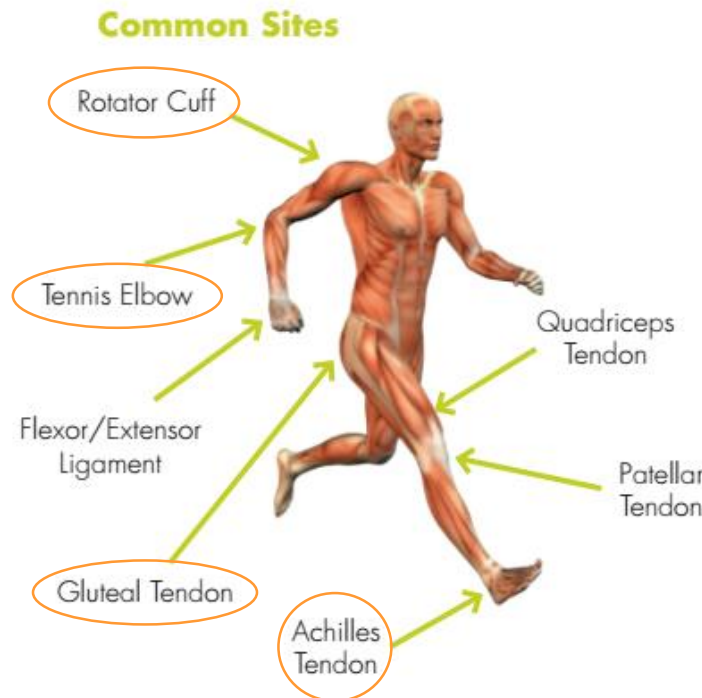
Orthocell's clinical development strategy is designed to increase the probability of achieving regulatory approvals, with significant levels of clinical validation achieved, and optimised manufacturing capabilities in place

Study ID	Description	No. of subjects	Treatment site	Duration (months)	Outcome Measures	Study status
CG-002	<i>Study of CelGro for dental guided bone regeneration around dental implants</i>	10	Dental	6	• Bone regeneration	Unpublished
CG-004	<i>Study of CelGro to augment surgical repair of rotator cuff tendinopathy and tear</i>	30	Shoulder	12	• Improvement in pain and function of shoulder • Quality of life • Measurement of tendon healing by MRI	Recruiting patients
CG-005	Treatment of hip cartilage defects with microfracture and CelGro	25	Hip	12	• Improvement in pain and function of hip • Return to sporting activities • Quality of life	Recruiting patients
CG-006	Surgical repair of peripheral nerve injury with CelGro	20	Upper limb	24	• Improvement in pain and function of arm/hand • Recovery of nerve function (sensory and motor) • Quality of life	Recruiting patients
CG-007	<i>Study of CelGro to augment arthroscopic repair of rotator cuff tendinopathy and tear</i>	30	Shoulder	12	• Improvement in pain and function of shoulder • Quality of life • Measurement of tendon healing by MRI	Awaiting ethical approval

What is tendinopathy?

- **Chronic degenerative tendon injury** usually brought on by repetitive motion
- Reduces healthy tenocyte population, tendon **cannot synthesise adequate collagen to restore normal structure**
- Disrupted tendon structure causes further injury, leading to a **cycle of injury, inadequate repair and degenerate tendon structure**
- Many people **cannot return to previous level of activity**

Where does it occur?



What causes it?

The severity of the injury is influenced by many factors, including:

- Amount of overuse
- Recovery time
- Genetic predisposition
- Ergonomics associated with the repetitive motion activity
- Age, level of fitness, general health
- Length of time before seeking help
- Quality of medical care

Ortho-ATI® - key advantages

Ortho-ATI® is the preferred therapy for degenerative tendon repair because it addresses the underlying pathology, is minimally invasive, cost effective with long term clinical data

Comparative advantages to traditional therapies

Traditional therapies only address the symptoms of tendinopathy (pain, immobility and stiffness), thus have limited long-term efficacy

- **Rest, physiotherapy and injections:** ineffective in the degenerative injury phase (more suited to acute injury)
- **PRP¹:** associated with symptomatic relief, but has inconsistent outcomes and duration of effect is not long-lasting
- **Corticosteroids:** reduces pain but weakens tendon structure long term (especially if repeated)
- **Surgery:** invasive, expensive, requires lengthy rehabilitation and increases economic burden on the healthcare system; patients cannot return to work / sport for months after treatment

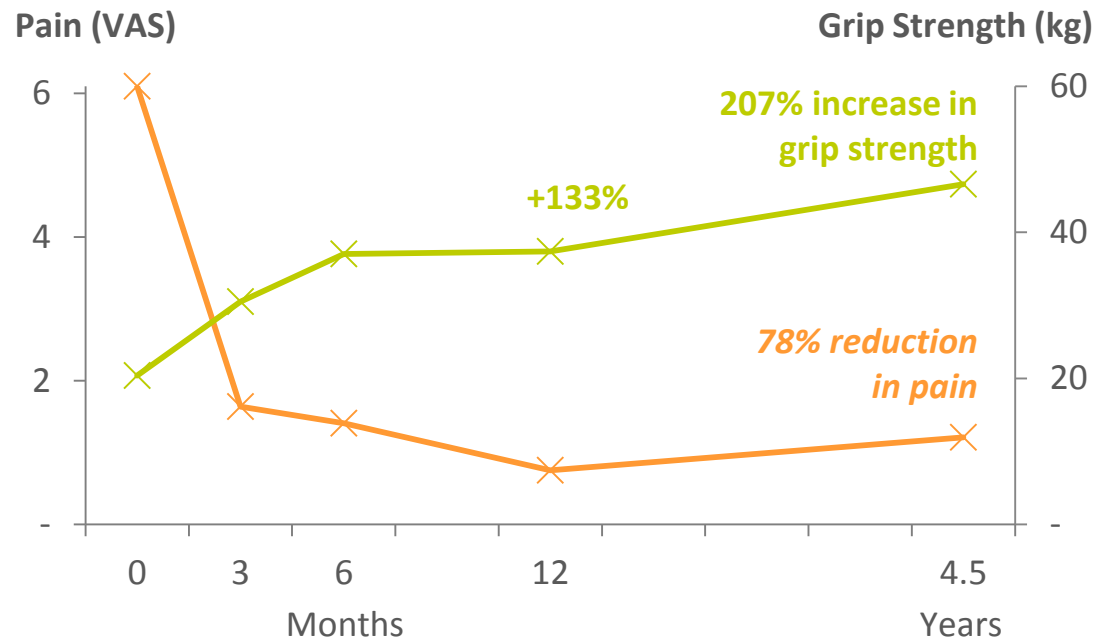
	Tenocytes (Ortho-ATI®)	PRP ¹	Corticosteroids	Surgery
Minimally invasive	✓	✓	✓	✗
Long term clinical data	✓	+/-	✗	+/-
Cost effective	✓	+/-	+/-	✗
Addresses underlying pathology	✓	✗	✗	✗

1. Platelet Rich Plasma Injection Therapy (PRP therapy)

Ortho-ATI® - clinical data

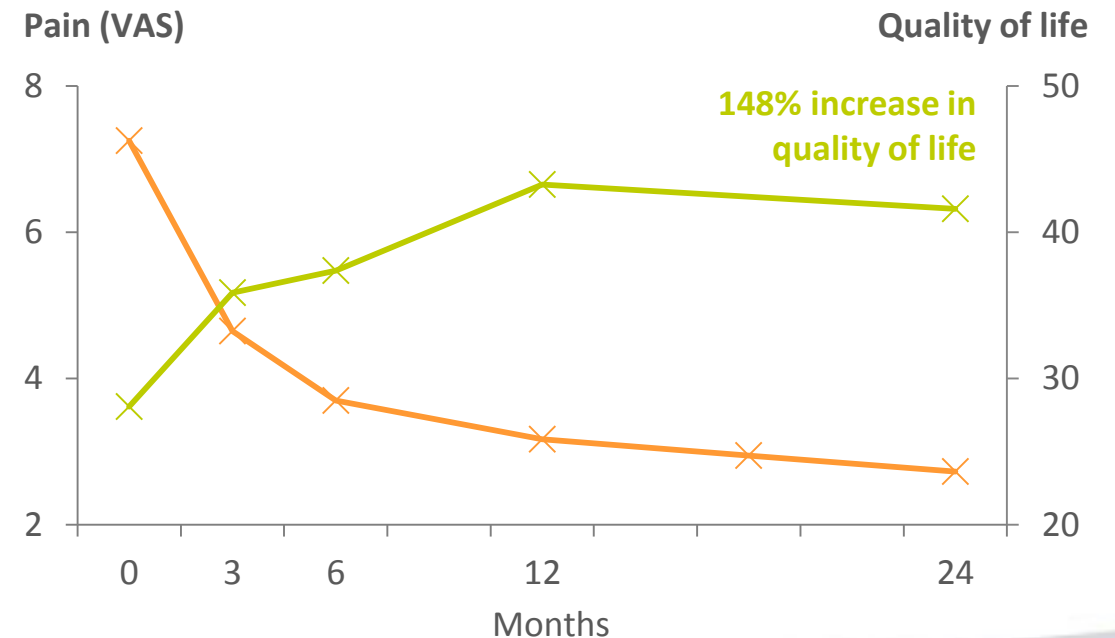
Ortho-ATI® tennis elbow study

- 17 patients that had failed all other treatments
- The patients had been experiencing symptoms for an average of 31 month
- Data to 4.5 years published in American Journal Sports Medicine



Ortho-ATI® gluteal tendon study

- 12 patients with gluteal tendinopathy – no remaining treatment options
- Average duration of symptoms of 33 months
- Significant clinical improvement in pain and function scores at 24 months



Ortho-ACI[®] and R&D pipeline

Orthocell is undertaking a “capital light” approach to commercialising Ortho-ACI[®] and developing other pipeline products which represent exciting potential value upside within the regenerative medicine space

Orthocell's foundation product

Ortho-ACI[®]

Marketable cartilage repair and regeneration therapy

- 3rd generation product
- >500 patient implants to date
- Included on the Australian Register of Therapeutic Goods, enabling the commencement of the process for reimbursement
- Cost effective treatment with potential for multiple applications



Ortho-ACI[®] procedure

Significant market opportunity

CelGro[®] pipeline

Potential for innovative platform technology solutions

- **ACL ligament replacement:** commence clinical study; positive results from initial animal study (recently completed)
- **Other potential applications:** General surgery, Urogynaecological, Collagen powder (for bone void fillers)



ACL ligament replacement

Significant commercial appeal expected in “off the shelf” products

Growth factors

Leverage existing knowledge towards human clinical trials

- Cell factory-derived tissue specific growth factors **to augment bone, cartilage and tendon repair**
- Part of successful **international collaboration**² – multiple animal studies completed and published
- “Cell Factory” **patents granted in key jurisdictions** (USA and Europe)

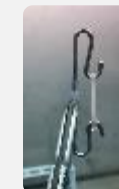


Growth factor samples

Lab grown tendons

Next generation product to complement Ortho-ATI[®]

- **Successfully manufactured** human tendon in a laboratory
- **Ongoing collaboration** with academic researchers¹ (received **ARC linkage grant**)
- Significant **unmet patient needs** and **multiple applications** (i.e. hand, shoulder and hamstring)



Lab grown tendon

1. Successful collaboration with Griffith University; University of Western Australia; La Trobe University; and University of Auckland
2. Successful collaboration with Lund University (Sweden); University of Western Australia; Indian Institute of Technology Kanpur (India)

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