

Tuesday 21 May 2024 – Draft programme

Pre-conference workshops			
9.00am – 5.00pm	ANDHealth Digital Summit The full-day event will be packed with engaging leaders and feature keynote presentations, panel discussions, and networking opportunities. Throughout the event, industry experts will share real world evidence and lived experience to address the challenges in developing, commercialising and implementing cutting-edge digital and connected health technologies, products and services into global healthcare systems. The Constitution of the digital Summit Incomplete Summit		
	The Summit will be attended by digital and connected health companies, industry professionals, service providers, investors and multinational medical device and pharmaceutical companies.		
8.30am – 12.30pm	ASEAN regulatory requirements - addressing the practical issues The ASEAN region is an important market for Australian medical device companies for a variety of reasons, such as its proximity (example being physically closer to Australia than Europe), large market size (population of 600+ million) and increasing demand for improved standard of care and therefore medical devices. ASEAN medical device regulations are guided by the ASEAN Medical Device Directive, introduced about a decade ago		
	In terms of regulatory approvals, ASEAN is different from European CE certification (CE Marking) process, with no version of CE Marking existing in the ASEAN region. Medical device companies need to seek regulatory approvals for each ASEAN country. The effort for medical device Regulatory Affairs team is not to be underestimated. The objectives of this workshop are to: Gain an appreciation of ASEAN medical device regulations and the implementation across each country. Get an update on current topics of interests on ASEAN medical device regulations. Clarify questions you might have in your preparation for registering medical devices in the region. (Please provide your questions in advance via email). Appreciation of the requirements for Good Distribution Practice Medical Devices (GDPMD). GDPMD has been and continues to be, rolled out across the ASEAN countries. GDPMD is a requirement for obtaining Import/wholesale/distribution license.		

1.00pm - 5.30pm

AusMedtechInvest 2024

AusMedtechInvest 2024 is Australia's inaugural boutique medtech investment roundtable forum, and aims to build meaningful personal connections between innovative medical technology (devices and diagnostic) businesses and investors, and help great ideas attract the capital needed to thrive in a fast-paced, competitive market.



1.30pm - 4.30pm

AusMedtech South Australian Site Tours







Adelaide Intermediary Program

Adelaide BioMed City



Tour One: Adelaide BioMed City
Tour time: 1:30 pm - 4:30 pm
Check-in: 1:00 pm - 1:30 pm

Meeting point: Adelaide Convention Centre - AusMedtech Registration Area

Transport: Walking tour

Tour Two: Lot Fourteen

Tour Time: 1.30 pm – 4.30pm

Check-in: 1:00 pm – 1:30 pm

Meeting point: Adelaide Convention Centre - AusMedtech Registration Area

Transport: Walking tour and tram

Tour Three: Tonsley Innovation District

Tour Time: 1.30 pm – 4.30pm **Check-in:** 1:00 pm – 1:30 pm

Meeting point: Adelaide Convention Centre - AusMedtech Registration Area

Transport: Coach to Tonsley and returning to the Adelaide Convention Centre. Refreshments will be provided.

Tour Four: Mawson Lakes | Australian National Fabrication Facility - South Australia

Tour Time: 1.30 pm – 4.30pm **Check-in:** 1:00 pm – 1:30 pm

Meeting point: Adelaide Convention Centre - AusMedtech Registration Area

Transport: Coach and walking tour. Refreshments will be provided.

5.30pm -7.00pm	Pre-Conference welcome reception
	AusMedtech 2024 Welcome reception
	Sponsored by:
	FB RICE
	The IP Navigators
7.30pm	Clinical trials dinner AusBiotech and its AusBiotech Clinical Trials Advisory Group warmly invite those working in the clinical trials sector to join them for this unsponsored, informal dinner and to connect face-to-face before AusMedtech 2024 conference officially begins.
	Clinical trials contribute to Australia economically and socially and are a critical component in the development process of bringing new therapies, devices and diagnostics to patients. However, the Australian clinical trial ecosystem is complex, involving many parts and stakeholders; this dinner is an opportunity to unite under the same roof and celebrate the sector and its progress.



Wednesday 22 May 2024 – Draft programme

07.30am –	Delegate registration	
08.30am		
8.30am –	Opening & welcome	
8.50am		
	Conference opening and welcome address	
	• The Hon Susan Close MP, Deputy Premier for South Australia, Minister for Industry, Innovation and Science, Minister for Climate, Environment and Water, and Minister	
	for Defence and Space Industries	
	Rosanne Hyland, Chief Operating Officer, AusBiotech	
8.50am-	Keynote session	
9.15am		
	National Health Medical Research Council (NHMRC) update	
	Chair: Gavin Fox-Smith, Chief Executive Officer, Omnigon Professor Steve Wesselingh, Chief Executive Officer, National Health Medical Research Council (NHMRC)	
	Further details to be confirmed	

9.15am –	Plenary session	
10.00am		
	Australian success stories	
	This panel discussion will celebrate the 2023/24 successful expansion and growth of medtech companies. Followed by a Q&A with the audience.	
	Chair: Robyn Lindner, General Manager, AusBioNSW	
	Panellists:	
	Sam Lanyon, Executive Director, Lumos	
	Marjan Mikel, Chief Executive Officer & Managing Director, Respiri	
	Paul Anderson, Chief Executive Officer & Managing Director, Orthocell	
	Jerneen Williams, Director of Operations, Bellberry Limited	
10.00am – 10.25am	Keynote session	
	Medical device regulation: an update on the latest	
	With sweeping reforms occurring across the globe in medical device regulation, this session focuses on how the Therapeutic Goods Administration (TGA) is supporting the industry	
	through these changes, along with the challenges of regulating new and emerging technologies and what this means for Australian patients.	
	Chair: Karen Parr, Director Policy & Communications, AusBiotech	
	Professor Anthony Lawler, Deputy Secretary, Medical Devices & Product Quality Division, Therapeutic Goods Administration	
10.25am –	Morning tea in the exhibition	
10.55am		
10.55am- 11.55am	Plenary session	
	Harnessing the pipeline: Partnering with big medtech	
	For medtech companies commercialising biomedical research across the ecosystem, there is an interdependent relationship between small and large companies. Multinational	
	companies are a critical part of the Australian landscape, with their ability to provide the resources, experience, and infrastructure necessary to support R&D, clinical development,	
	manufacturing, and distribution of devices and diagnostics.	
	This plenary session provides an invaluable opportunity to hear from big medtech on partnering, prospecting, and support work in the Australian environment.	
	Chair: Dell Kingsford Smith, AusBiotech Board; Vice President Medical Affairs, Market Access & Government Affairs, Asia Pacific, Cochlear	
	Panellists:	
	Maurice Ben-Mayor, President & Managing Director, Stryker	
	Mick Trevaskis, Chief Executive Officer, Device Technologies	
	Rebecca Cortiula, Senior Managing Director Australasia, Varian	
	Jane Crowe, Managing Director, ANZ Cardinal Health	
	Pat Williams, Vice President & Country Manager ANZ & Korea, Edwards Lifesciences	
11.55am –	Room move break	
12.00pm		

How to navigate MDR, why is the EU different now?		
Europe continues to face a dynamic and challenging regulatory environment. In effect since 26 May 2021, the European Union's (EU) medical device (and In-Vitro Device Regulation (IVDR)) regulations mandate that medical device manufacturers targeting the EU market must adhere to new standards. Key changes introduced by these regulations include: New requirements for translations Stricter clinical evaluation and post-market surveillance Expanded product scope Introduction of unique device identification (UDI) systems Greater transparency and data reporting Reinforced patient safety measures, and Enhanced role of notified bodies This panel of seasoned experts will discuss the issues facing the sector and pitfalls new players often come across. Chair: George Loizou, Director, CMS SciDoc Pty Ltd Panellists: Julie Winson, Quality and Regulatory Director, LBT Innovations Anthony Skeats, Chief Operating Officer, Micro-X Chris Henry, Managing Director, Actis Medical Hwee EE Tan, Principal Consultant, DH RegSys Consulting Pty Ltd	Building workforce skills and capabilities for successful medical device commercialisation In this session facilitated by ARCS Australia, the focus is on workforce skills and capabilities crucial for the effective commercialisation of medical devices. The session will feature representatives from industry, academia, and government perspectives to provide cross sector insights. Facilitated discussions will emphasise cross-sector collaborations, aiming to empower attendees with actionable insights for navigating the intricate landscape of medical device commercialisation. This session aspires to foster collaboration and skill enhancement, ensuring successful market entry for medical innovations. Chair: Dr Tim Boyle, Chief Executive Officer, ARCS Panellists: Dana Bell, Director Partnership SA, MTPConnect Shan-Shan Wang, Founder & Chief Executive Officer, Roam Technologies (RoamTech.ai) Ajay Nair, President APAC, Mullings Group Paul Cohen, Managing Director & Founder, Paul Cohen Consulting In partnership with:	After almost 2 years of steep declines in valuation, many industry players are talking about a "return to normal". But were the heady days of no-low due diligence, FOMO driven investments and "frothy" valuations really normality? Or are we just longing for a moment in time to return? Chair: Bronwyn Le Grice, Managing Director & Chief Executive Officer, ANDHealth Panellists: Dr Chris Nave, Founding Partner & Managing Director Brandon Capital Dr Amanda Gillon, Senior Partner, BioScience Managers Dr Steve Burnell, Managing Director, Ten Mile Elyse Shapiro, Healthcare Analyst, Canaccord Genuity

Concurrent	Concurrent sessions		
1.55pm	Session D	Session E	Early-Stage Innovation Forum
2.50pm	Session D	Session E	Larry-Stage Illilovation Forum
2.50pm	Ethics and AI – has the horse already left the stable?	Demystifying federal funding programs	The Early-Stage Innovation Forum, will feature rapid fire rounds of quick-pitch presentations on early-stage technologies and projects
	Artificial intelligence has been heralded as the largest technological change facing our generation, but concerns about ethics, privacy and consumer protection are still front of mind. As the Australian Government contemplates regulation of AI, we have to consider whether this might be a case of shutting the stable door after the horse has bolted. Chair: Grace Lethlean, Chief Product Officer, ANDHealth	Set yourself up for funding success by learning about the suite of programmes funded by the Federal Government that support medtech innovation. Hear about the purpose and intention of each programme, eligibility criteria, who they are best suited for, the type of projects they are looking to support, and upcoming dates. Chair: Dr Tracey Wilkinson, Director Stakeholder Engagement WA, MTPConnect	from local research institutes, universities, hospitals and pre-series A companies in the areas of medical devices and diagnostics, digital health and enabling technologies.
	 Panellists: Daniela Cecere-Palazzo, Senior Lawyer, Youlegal Angie Corbo, Informatics Global Catalyst Innovation & Architecture, Roche Liesl Yearsley, Chief Executive Officer & Founder, Akin Justine Lacey, Director, Responsible Innovation Future Science Platform, CSIRO In partnership with: 	 Panellists: Associate Professor Tracey Laba, Director Frontiers Program, Medical Research Future Fund, Dept of Health & Aged Care Representative, National Reconstruction Fund David Chuter, Executive Director, Industry Growth, Department of Industry, Science and Resources Liz Crompton, SME Connect Program Advisor, CSIRO 	
2.50pm- 3.45pm	Session F	Session G	Early-Stage Innovation Forum
	Making market access a priority for Australian Innovators In 2017 the OECD released a report detailing that "public procurement offers an enormous potential market for innovative products and services. Used strategically, it can help governments boost	The growing importance of environmental, social, and governance (ESG) and how does the medtech board react to this? The Boston Consulting Group states "Health care accounts for 5% of total global carbon emissions, and	
	innovation at both the national and local level and ultimately improve productivity and inclusiveness." In	medical devices and technology are responsible for a large portion of that. Much of this comes from the manufacturing operations and supply chains of medtech	

Australia it could be argued that this has not been the case and the Australian response to the COVID19 pandemic is emblematic of this attitude in our healthcare system in particular. Issues that could be driving this attitude could include such factors as risk aversion, management, personnel and skills capacity and political support.

This experienced panel will discuss possible solutions such as policy strategies with defined targets, sufficient budgets, funds and other financial incentives, build staff capabilities and skills, setting up multidisciplinary teams and competence centres focused on public procurement for innovation.

Chair: Tracey Shearer, Managing Director, August Consulting

Panellists:

- Professor Tracy Merlin, Deputy Executive
 Dean, Faculty of Health and Medical Sciences
 Head, School of Public Health, University of
 Adelaide
- Andrea Andrews, Executive Director
 Procurement and Supply Chain Management,
 SA Health
- Gibran Maher, Founder and CEO, Additive Surgical
- Hon Stephen Dawson MLC, Minister for Emergency Services; Innovation and the Digital Economy; Science; Medical Research; Minister Assisting the Minister for State and Industry Development, Jobs and Trade

companies and their suppliers. At the provider level, medtech generates tonnes of unrecycled waste through single-use disposable products and packaging."
There is a growing demand from customers, shareholders and employees for Boards to implement ESG initiatives within organisations. Medtech companies must choose to balance current competitiveness and future investments while connecting ESG to economic value. This panel of experienced executives and Board members will tackle the thorny issue of how to do this whilst generating value.

Chair: Melissa McBurnie, Partner & Head of Impact, Brandon Capital

Panellists:

- Olivia Pitt, Head of ESG, Ellerston Capital
- **Lis Boyce**, Partner, Piper Alderman
- Elizabeth Dallimore, Executive Chair, Inspiring Holdings
- Dr David Brookes, Executive Chair, Anatara Lifesciences

3.45pm-4.15pm

Afternoon tea in the exhibition

Concurrent sessions

4.15pm- Session H Session I Early-Stage Innovation Forum
5.10pm

Beyond patents: strategic use of all your intangible Reimbursement assets and how not to kill a deal Are Australia's medtech reimbursement pathways appropriately designed and sufficiently rewarding to Our panel will discuss the often-untapped value that sits attract both local and global innovation? within a corporate's intellectual property (IP) and provide tips to identify, grow, and maximise the value of Chair: Dell Kingsford Smith, VP Medical Affairs, these assets. As well as a focus on building one's own IP market access and Government Affairs, Cochlear and market exclusivity, the panel will also discuss the importance of understanding the IP held by others and the resultant freedom to operate in key markets. **Panellists:** • Penny Shakespeare, Deputy Secretary, Chair: Rachel Hooke, Partner, FB Rice Department of Health and Aged Care • Polo Guilbert – Wright, Senior Director Panellists: **Anna Smyth**, Partner, Gilbert and Tobin Government Affairs ANZ, Edwards Derek Minihane, Partner, Deloitte Lifesciences John Heasman, Chief Operating Officer, George Papadopoulos, Director & Partner, **Epiminder** Lucid Healthcare Consulting • Sarah Griffin, Director, MedTechnique Consulting Nicola Leavold, Commercial Director Australasia, BXTA Pre-dinner drinks 7.00pm-7.30pm 7.30pm -AusMedtech conference dinner 10.00pm Rosanne Hyland, Chief Operating Officer, AusBiotech Dinner speakers: Scott Stirling, Chief Executive Officer, Red Dust, and Jonathon Lindsay-Tjapaltjarri Hermawan, Director of Programs and Strategy Lead, Red Dust Facilitator: Rebekah Cassidy, Chief Executive Officer, AusBiotech With Australia's indigenous communities overrepresented in low socioeconomic figures, Australia's First Nations people face a greater risk of poor health, higher rates of illness, disability, and death, and live shorter lives than people from higher socioeconomic groups. Founded in the Northern Territory, not-for-profit Red Dust draws on the strengths of both western health knowledge and traditional cultural knowledges to create a positive influence on young people and improve outcomes for communities. Providing a 'community-as-family' model of health and well-being programmes, Red Dust works alongside community leaders and elders to create a stronger future for indigenous youth and their families. Join us for a fireside chat with Red Dust founders, Pintupi man Jonathan Hermawan and CEO Scott Stirling, as they discuss their personal journey over the past seven years growing Red Dust and developing a cooperative model that is charting a new course and leading to improved health and wellbeing outcomes.



Thursday 23 May 2024 – Draft programme

8.30am –	Delegate registration
9.00am	
9.00am –	Hon Stephen Dawson MLC, Minister for Emergency Services; Innovation and the Digital Economy; Science; Medical Research; Minister Assisting the Minister for State and Industry
9.15am	Development, Jobs and Trade
9.15am –	Plenary session
10.00am	
	Issues facing healthcare in Australia Get your caffeine fix from the exhibition hall and start day 2 of the conference with a wide-ranging panel discussion with leaders from our representative industry and government organisations about the burning challenges- and the exciting opportunities- facing the Australian healthcare sector that are keeping them awake at night. Chair: Stuart Dignam, Chief Executive Officer, MTPConnect
	Panellists: Ian Burgess, Chief Executive Officer, MTAA Dr Anna Lavelle, Chair, Medicines Australia

	 Dr Tim Boyle, Chief Executive Officer, ARCS Bronwyn Le Grice, Managing Director & Chief Exe Rebekah Cassidy, Chief Executive Officer, AusBiote 	•	
10.00am – 10.45am	Keynote session		
	receiving end of verbal and physical harassment and abuse media, and private industry, with insults, abuse, threats, tro	reflect on their personal experiences as they shine a spotlig nation against women in the workplace. tive Officer, ANDHealth 8 & NED Hansen Technologies	t occurrence for women in politics and advocacy, academia,
10.45am – 11.15am	Morning tea in the exhibition		
11.15am – 12.10pm	Session J	Session K	Session L
	Al in medical imaging – from promise to practice	Exploring the emerging material science manufacturing and investment opportunities to reduce medical plastic	Optimising clinical trial delivery
	The promise of artificial intelligence (AI) in medical imaging analysis has been discussed for many years and is now a reality. Was it smooth sailing for this seemingly ideal use case? Are there lessons to be shared which will inform the next wave of technologies? In this panel discussion we'll hear from those who have been at the forefront of the development and adoption of AI in medical imaging, they'll share what's worked, what didn't, the challenges that remain and where to next. Chair: Dr Ludovic Labat, Chief Executive Officer, Neo-Bionica Panellists: • Dr Michelle Perugini, Chief Executive Officer, Presagen Dr Mark Phillips, Head of Clinical Research & Medical Affairs, Annalise.ai	Waste Hospital generated plastic waste is a worldwide issue. When first introduced into hospitals, single-use plastics were an attractive option as it allowed for maintenance of a sterile environment and infected plastic material could be easily disposed via landfill waste. For equipment such as syringes this is vital. However, the sheer quantity of single use plastics being used in hospitals is becoming alarming, particularly considering the overuse or unnecessary use of single-use plastics. In Australia, the healthcare industry is responsible for around 8% of Australia's carbon emissions and generates large amounts of non-recyclable plastic waste. Environmental Social Governance (ESG) factors are increasingly becoming a factor in procurement and related commercial decisions.	As an early stage medtech company or founder it is often difficult to know exactly what is required to get a product/asset through the clinical phase. This session will explore the things you need to consider when running a clinical trial and before you start writing the protocol. You have a product, you know what you want it to do, you've even got it to the point it's ready for clinical testing and met all the manufacturing regulatory requirements. But wait, clinical research has its own regulatory standards and requirements. How on earth do I navigate this world? The panel will discuss the considerations for embarking on the clinical development phase of a trial, when you should start to plan, what to look out for, whether to hire or contract expertise and how to engage a site(s) and clinician(s). Chair: Natalie Barber, Director, Clinical Operations, Chrysalis

	 Dr Wenji Pang, Chief Scientific Officer - Imaging & Al, Resonance Health Alison Deslandes, President, Australasian Society of Ultrasound in Medicine 	This diverse panel session will discuss ways to address these issues and funding assistance that is available. Chair: Stuart Anderson, Clinical Translation and Commercialisation Medtech Program Manager, MTAA Panellists: Jane Crowe, Managing Director, ANZ Cardinal Health The Padam Walczak, Director, HNE Innovation Office, Hunter New England Local Health District Pauline Salib, Antimicrobial Stewardship Pharmacist, Western Health Leonie Walsh, Interim Chair, Solving Plastic Waste CRC The Carly Hollier, A/Senior Manager, Sustainability, Delivery Excellence, Partnerships & Projects, HealthShare NSW In partnership with:	 Simon Cook, Executive Director, Operations, Eudaemon Technologies Helen Plummer, Research Manager, Cercare Prof Les Bokey, Institute Director, Ingham Institute for Applied Medical Research Simon Belcher, General Manager, Three Peaks Medical
12.10pm – 1.05pm	Session M	Session N	Session O
·	Strategic considerations in manufacturing for medtech With a strong push and incentives for sovereign and local manufacturing capacity, how might medtech companies think about whether to buy, build or partner to develop the capabilities, skillsets and tools required to execute on manufacturing at scale. This session will explore the strategic thinking required for manufacturing business decisions, including the impact of technologies such as additive manufacturing, Industry 4.0, robotics, AI and other emerging technologies.	What angel investors look for in a MedTech startup company Many early-stage companies require external funding. One source of funding can come from Angel Investors, who typically provide a cash injection in exchange for equity in the company. Startups, particularly in the medtech sector, can be a risky proposition for an investor, and there are key elements that an Angel Investor will look for when deciding whether to invest in that company. Such considerations may include the strength of the team behind the startup, the business	Are medtech and digital product clinical trials - same or different? Following on from the previous panel session that explored how early stage medtech companies navigate the clinical phase, in this session the invited speakers explore more specifically the commonalities and differences in designing, running and managing clinical trials between digital products and devices, as well as digitally enabled devices. Founder will share how they navigated the sometimes-unclear requirements for Software as a Medical Device trials

model, what intellectual property the startup has, and

ultimately, what sort of return the Angel Investor can

obtain on their investment.

Advisory

Chair: Nick Northcott, Managing Partner, Chrysalis

in both Australia and the US, and what they learned from

how to work with CROs, and how to assess who would be

the more established medical device trials. We also explore

Panellists: the right partner. This panel session, which comprises angelilnvestors who The goal of the session is to get clarity on what common Lisa Henretty, Chief Operating Officer, Enersol pitfalls are, and highlight relevant factors and strategies to Louisa De Vries, Consulting Manager, Bosch are active in investing in startups in the medeech industry, will discuss the key elements that they look for accelerate the clinical phase for both devices and digital **Australia Manufacturing Solutions** in a startup when deciding whether or not to invest, and products. Val Valentine, Director, Edwards Lifesciences provide insight into how you, as a startup, can best Pablo Solis. Chief Executive Officer & Coposition yourself so as to make yourself as attractive to Chair: Dr Katja Beitat, Head of Health Tech, Cicada Founder, Protego Medical an angel investor as possible. Some of the panel Innovations members have also been in the position of forming a Panellists: startup and themselves seeking investment, and can provide valuable advice from their own experiences. **Helen Souris**, Chief Executive Officer, CardiHub The session will be chaired by Dr Milena Dryza, Senior Mary-Beth Brinson, Chief Executive Officer Cyban Associate of Madderns Patent and Trade Mark Stewart Bartlett, Chief Executive Officer Ferronova Attorneys. Milena is a patent attorney with many years Eric Davies, Former VP of Global Marketing for of experience in private practice, as well as an in-house Abbott and Cochlear, advisor to Roam patent counsel for leading biotechnology company, CSL. Technologies (RoamTech.ai) Chair: Dr Milena Dryza Senior Associate, Madderns Patent & Trademark Attorneys. Panellists: Dr Anabela Correia, Chief Executive Officer, LiVac **Dr Nick Haan,** Chief Executive Officer. Seonix **David Saint**, Chair, Southern Angels Dr Amandeep Hansra, Lead Investor and Cofounder of Medical Angels Sponsored by: madderns 1.05pm -Session P Session O Session R 2.00pm What infrastructure do we need to build a vibrant How to close a VC round (and retain your sanity) Partnering - the pathway to success...or is it? medtech innovation and commercialisation ecosystem Competition for capital is fierce at the moment, with A business partnership is a collaboration between two or investors focusing on the stability of their existing more entities that pool resources, technology and/or The road to successful commercialisation in medtech is portfolio and wary of cash burn in a tight market. So finances to achieve a generally agreed goal. In medtech this often winding and with many obstacles along the way.

Being aware of the pitfalls and shaping your start-up for the best possible outcome is no simple task. It has been said we need the right people, science, infrastructure and money to build a success. Incubator and accelerator programmes are a vital part of the medtech innovation ecosystem but what else is required to help our sector? and provide a rich environment of education, mentorship, industry collaboration, networking and capital raising support. Our panel will look into the roles of education, mentorship, industry collaboration, networking and capital raising support. Where are the gaps and how do we fill them?

Chair: Professor Karen Reynolds, Director, Medical Device Partnering Program

Panellists:

Lunch in the exhibition

2.05pm -

- Dr Lilly Bojarski, General Manager, Cicada Health Tech Hub
- Natalie Rickers, Commercialisation Director, South Australia MTP Connect
- Kelly Coverdale, Managing Director, Cover Biomedical
- Professor Sharath Sriram, Director, Discovery to Device Facility, RMIT University; President, Science & Technology Australia

how DO you land a significant investor, and more importantly, what do you need to do to close the deal!

Chair: Sarah Meibusch, Partner, OneVentures

Panellists:

- Arthur Shih, Chief Executive Officer, Humanetix
- Peter Vranes, Chief Executive Officer & Co-Founder, Nutromics
- Richard Horton, Partner, Squire Patton Boggs
- Dr Melissa McBurnie, Partner & Head of Impact, Brandon Capital.

is a widely accepted pathway to develop technology, access markets or ensure access to specific technologies. Whilst there are many instances of this being a success there are also many instances where the expected success did not materialise. This panel of seasoned executives will discuss many of the pitfalls and key decision points where partnerships are useful of inhibitory.

Chair: Peter Bradley, Principal, Qatalyst Consulting

Panellists:

- Peter Rowland, Non-Executive Director, Micro-X
- Michael Kavanagh, Chief Executive Officer & President, Nanosonics
- **Dr Anabela Correia,** Chief Executive Officer, LiVac
- Further speakers to be confirmed

3.00pm	Lunch in the exhibition
3.00pm –	Plenary session
3.30pm	
	Lessons from 20 years of collaborations creating meaningful medtech ventures
	With a global presence and multidisciplinary skills, IDE Group is a well-regarded partner for medtech commercialisation and product development.
	IDE works with its partners to find and assess business opportunities, conduct research, create and implement commercial strategy, gain access to funding, develop new technology, manufacture quality products and create successful medtech ventures. Since 2003, IDE has grown over 150 medical technology businesses and realised over 500 projects across the medical technology landscape globally.
	You will hear from IDE Group CEO George Sidis and Eudaemon Technologies Executive Director, Operations, Simon Cook as they discuss the organisation's journey to date and how it's supported Australian medtech innovators to generate high-impact ventures and improve health outcomes. Chair: Warren Bingham, Global Vice President, ARIA Research

George Sidis, Chief Executive Officer, IDE Group & Simon Cook, Executive Director, Operations, Eudaemon Technologies

3.30pm – 4.15pm	Plenary session
	Take home lessons from AusMedtech 2024 The two days of AusMedtech are a cornucopia of data, information, knowledge, and wisdom. Many of us will have individual gems that are pertinent to our particular business or problem to take home. However, many of the big picture and relevant issues explored may be missed because of conflicts or competing priorities drawing attendees to other sessions. This panel of hardened Medtech executives have made it a mission to pick the pearls of wisdom from the conference and highlight to the final audience. We will also draw on the collective attendees to reduce these to actionable take home messages from the conference. Chair: Peter Bradley, Principal, Qatalyst Consulting Panellists: Jo Close, Director Adelaide Intermediary Program, MTPConnect Craig Newton, Chief Executive Officer, Kynetyka
4.15pm –	- Further speakers to be confirmed Conference closing reception
5.00pm	Exhibition Hall
	Rosanne Hyland, Chief Operating Officer, AusBiotech