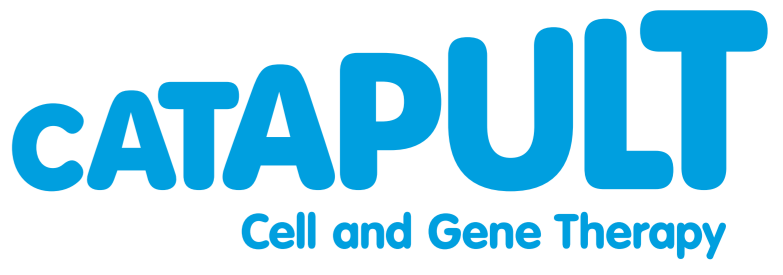
Towards Industry 4.0: Development of a Smart and Advanced Process Control for Autologous Cell Therapy CATAPUL P Statham 1, D Marginean 1, N Thompson 1, M Filipiak 1, S Wadud 1. C Pararasa 1 C Norinov 1 101 1 77 Bioprocessing Platform Integrating Real-Time Monitoring

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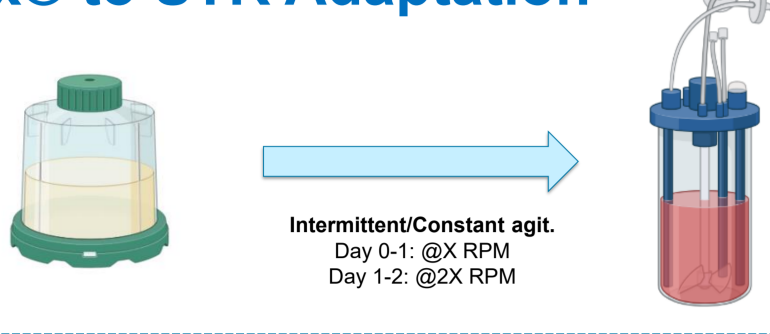


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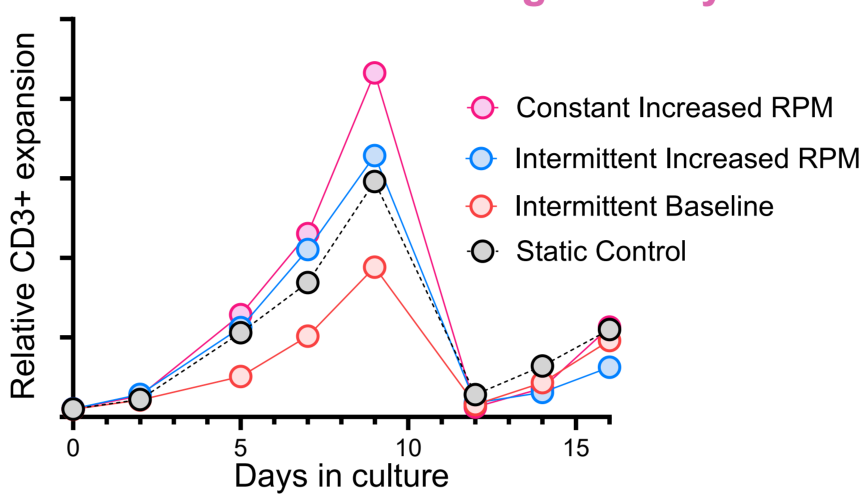
Clinical adoption and patient's accessibility of autologous immunotherapies remain hindered by the lack of efficient, consistent, cost-effective, and scalable manufacturing processes. To address this bottleneck, an international EU-funded consortium is developing a first-in-class, smart bioprocessing platform for personalised autologous cell therapies, integrating in-line process analytical technologies and advanced process control systems. Using a clinical-stage tumorinfiltrating lymphocytes (TILs) manufacturing process as a model, a static baseline process in the G-Rex system was adapted to a dynamic, stirred-tank reactor (STR) system. In addition, metabolomic profiling of spent media identified ~50 metabolites with a significant effect on cell expansion. Focusing on three key metabolites, alongside agitation speed and dissolved oxygen as critical process parameters (CPPs), we conducted design of experiment (DoE) studies using the STR systems, with integration of Raman probes, as an online process analytical tool. Based on these results, subsequent work will implement real-time monitoring and model predictive control strategies leveraging chemometric and mechanistic models to enhance process consistency and performance.

GRex® to STR Adaptation

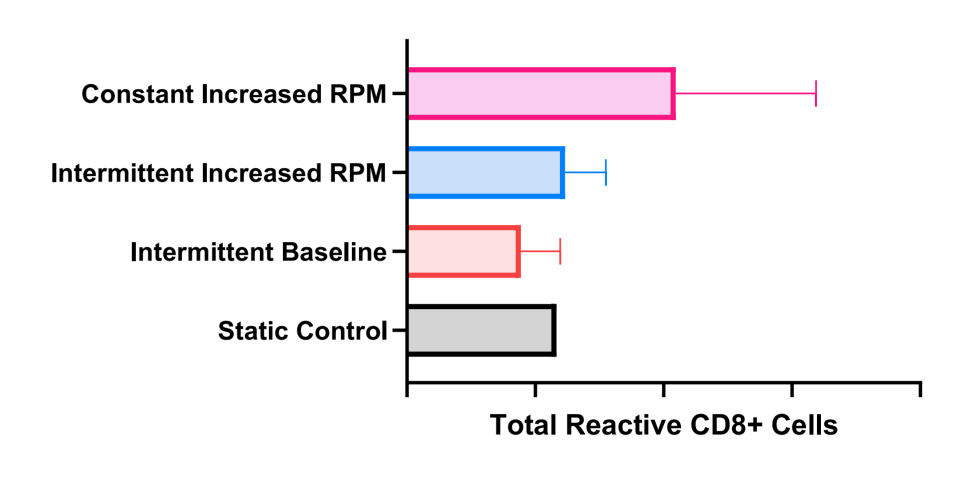


- Exemplar autologous TIL-DC co-culture process with subsequent reactive TIL expansion.
- Prior art suggest static conditions were thought to be needed for higher reactivities.
- Agitated systems are controllable. pH, DO, temperature, metabolite controls can facilitate improved autologous process performance.

Successful Translation to Agitated System

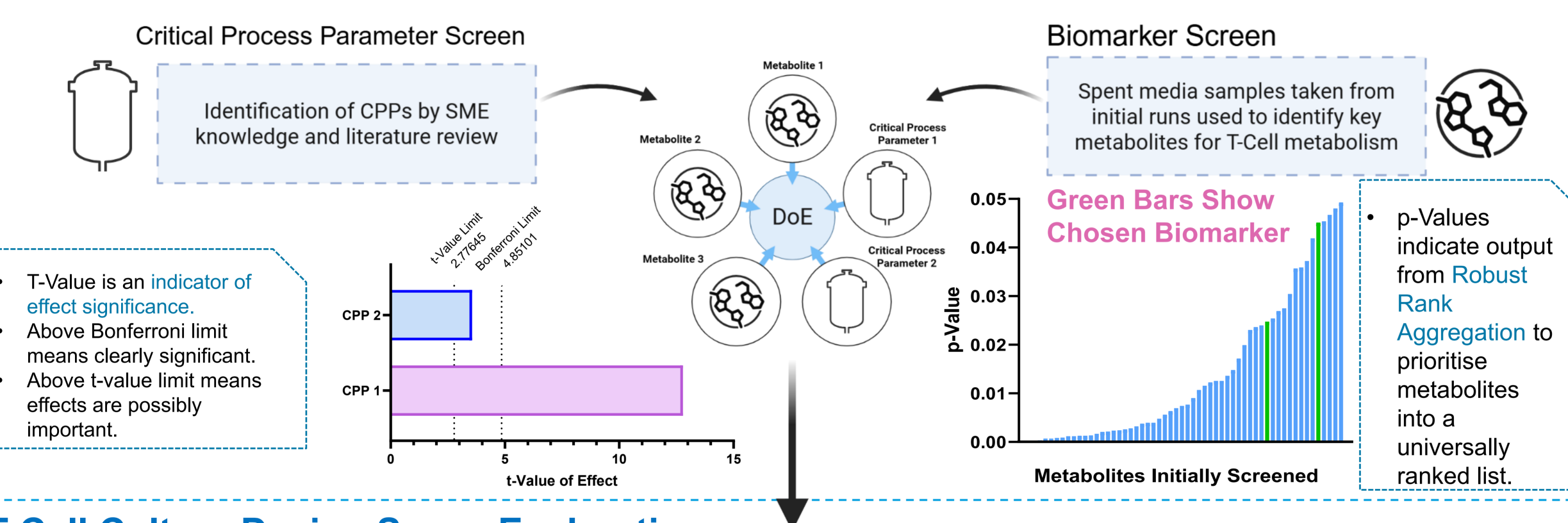


Maintenance of CD8+ Reactivity in Agitated System



- Successful translation of static process to STR and subsequent investigation of intermittent versus constant agitation.
- These studies showed constant, high RPM led to higher T-Cell expansion and reactivity than the initial static.

Critical Process Parameter/Biomarker Screen



- Two steps done in parallel.
- 7 CPPs were screened over 2 screening runs.
 - These resulted in identifying 2 process parameters critical to T-Cell expansion and reactivity.
- Biomarker screen led a shortlist of 57 metabolites.
- These were narrowed to 3 identifiable by Raman, and interesting for T-Cell metabolism.

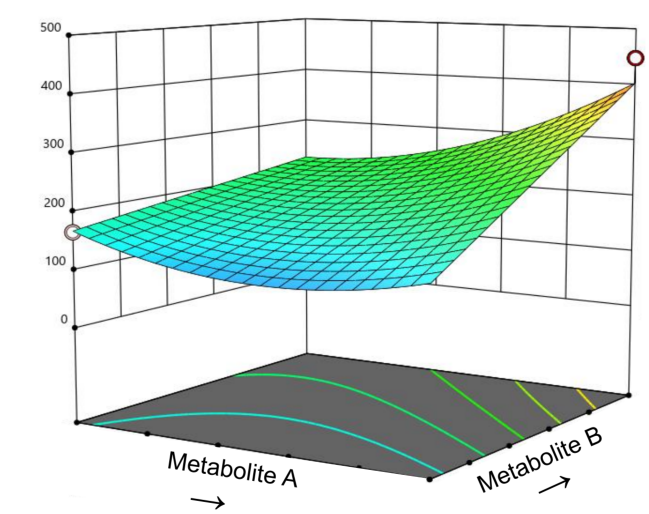
T-Cell Culture Design Space Exploration

- The DoE was a 5-level, central composite design, consisting of 32 STR vessels utilising parameters identified in the previous screens.
- Aim was to introduce significant process perturbations and measure their effects expansion of CD3+, reactive CD4+, and reactive CD8+.
- Raman spectroscopy data was collected for generation of a PAT chemometric model.

High CD3+ Fold-Expansion 500-400 Metabolite A (↓↓) Metabolite A (→) Metabolite B (↓↓ Metabolite B (\rightarrow) Metabolite B (\rightarrow) Metabolite C (↓) CPP 1 (↑), CPP 2 (↓) CPP 1 ($\downarrow\downarrow$), CPP 2 (\rightarrow)

- Conditions were used to elicit both positive and negative responses.
- Large range of responses allows for more robust bioprocess and chemometric models.
- Yield and reactivity improvements indicated optimal process conditions leading to 450-fold expansion.
- Perturbations allowed coverage of a large design space.

DoE Model Generation

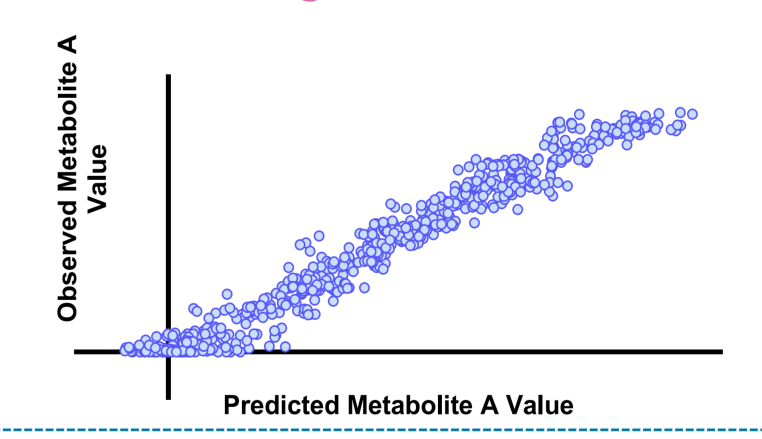


Optimal Parameter Determination

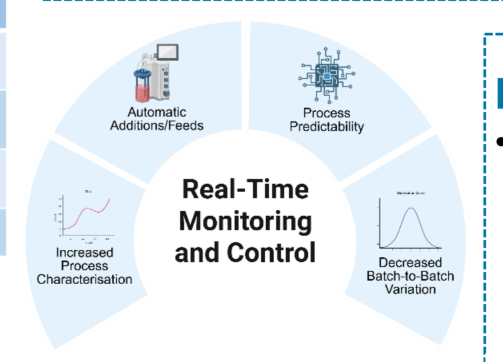
Metabolite A	Metabolite B	Metabolite C	CPP 1	CPP 2	Response	Response Prediction	Desirability
1	1	\	↑	\	↑ CD3+ Fold Expansion	403-fold	0.653
\	1	\	1	1	↑ CD4+ Fold Expansion	469-fold	0.224
\	1	\	↑	1	↑ CD8+ Fold Expansion	986-fold	0.696
\downarrow	\downarrow	↑	↑	↑	↑ % Reactivity	NA	0.462

- DoE model predicted ideal operating conditions for T-Cell bioprocessing.
- Model predicts that a combination of high metabolite A and high metabolite B will result in high CD3+ fold-expansion.
- Model also predicts that high metabolite C will increase % reactivity.

Generation of High R² Chemometric Models



- 800 Raman spectra were selected as training data for chemometric models.
- These models will be utilised for a model for predictive control as a strategy to predict process output based on real-time monitoring of CPPs.



Future Work:

- Generate and integrate chemometric and bioprocess models for use in real-time monitoring and control with healthy and patient material.
- Models will be tested for real-time monitoring of the CPPs and metabolites in future runs.
- Readings will inform a control system to automatically supplement feeds with metabolites to maintain an optimal concentration throughout the process.





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Schematics created using BioRender **Cell and Gene Therapy Catapult**

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