Aptamers 2017 at Oxford

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ABSTRACT

The 4th annual symposium of the International Society on Aptamers, Aptamers 2017, was held in Oxford on the 11th and 12th April and was well attended, with presenters from Europe (Spain, Germany, Austria, and UK), North and South America, Asia, and Australasia presenting on a diverse range of topics, from enhancing SELEX, to diagnostic applications such as lateral flow devices or medical imaging, to therapeutic applications such as drug delivery. The conference was split into six sections in total, covering chemical modifications, disease, analysis/diagnostics, tools/selection/design, riboswitches, and computation with over 50 oral and poster presentations. The conference started with a welcome from both the Symposium Chair, Professor Dr Ulrich Hahn (University of Hamburg, Germany) and the President of the International Society on Aptamers, Dr Sarah Shigdar (Deakin University, Australia).

CHEMICAL MODIFICATIONS OF APTAMERS

Aptamers are developing for a number of applications and they have demonstrated high potential for therapeutic applications. However, some aptamers are prone to rapid nuclease degradation. Therefore, in order to develop aptamers for clinical applications, some modification of the nucleotides are required (Shigdar et al, 2013). Professor Philip Johnson (York University, Canada) chaired this first session where we heard from a number of leading experts in the field. Professor Jesper Wengel (University of Southern Denmark, Denmark) co-inventor of locked nucleic acids, started off the first session by providing an update on the use of Phusion™ High Fidelity DNA polymerase and engineered polymerases as a way forward for SELEX using LNAs (Lou et al, 2017b), definitely something that has potential for better bioavailability of aptamers in therapeutic applications. Professor Wengel also discussed the use of LNAs and unlocked nucleic acids as antisense or RNAi drug candidates before presenting results on how to build artificial protein mimics through the combination of oligonucleotides and short peptide sequences (Lou et al, 2017a). Dr Xianbin Yang (AM Biotechnologies, USA) from AM Biotechnologies, then presented information on X-aptamers which are developed using microbead based single-cycle discovery process and does not rely on PCR amplification, allowing a variety of chemical modifications to be incorporated (Lokesh et al, 2017). Dr Dan Schneider (SomaLogic, USA) from SomaLogic was the third presenter in this session and he talked about enhancing chemical diversity with multiple
pyrimidine modifications. These modifications can create novel intramolecular motifs leading to improved aptamer affinities (Gawande et al, 2017).

**DISEASE**

One of the areas in which aptamers are showing a lot of potential is the area of theranostics. Aptamers bind to their target through complex interactions involving hydrogen bonding, van der Waals forces, hydrophobic, and electrostatic interactions. In this manner, aptamers can act as agonists or antagonists, or through receptor mediated endocytosis, the aptamers can be used for targeted drug delivery (Macdonald et al, 2018). Additionally, aptamers can demonstrate higher sensitivity in diagnostic assays, especially in the case of complex diseases. The second session, chaired by Professor Dr Beatrix Suess (Technical University Darmstadt, Germany), discussed various different ways aptamers are being used to effectively target and treat, or more sensitively diagnose disease. Dr Greg Penner (NeoNeuro SAS, Paris, France) started the session with a discussion of aptamer theranostics for Alzheimer’s disease. Dr Penner’s team preselected the randomised library by injecting the library into the tail vein of mice and then collecting only the aptamers that has passed into the brain. They then performed aptamer selection on human blood from Alzheimer’s patients to pick out targets that could be used as a diagnostic panel. Their panel was effective in stratifying patients into early, middle and late stage Alzheimer’s disease and they are now investigating this Aptamarker AD fingerprint for earlier diagnosis of disease (Lecocq et al, 2018). Next was Professor Dr Ulrich Hahn (University of Hamburg, Germany) and our Symposium Chair, who presented data on selectin and integrin aptamers as a means of preventing metastasis. Both of these aptamers were shown to have an anti-adhesive effect on cancer cells though truncation of the integrin binding aptamer had a detrimental effect on the function of the aptamer (Berg et al, 2016). Professor Yoshikazu Nakamura (Ribomic Inc, Japan) presented on a fibrolast growth factor 2 (FGF-2) binding aptamer that can restore bone growth in Acondroplasia (ACH) and is now gearing up for clinical trials. This aptamer has restored body weight and length in ACH transgenic mice and has an excellent profile in rat and monkey plasma. They are now looking at it in age-related macular degeneration (Jin et al, 2016). Professor Fernando Pastor (Foundation for Applied Medical Research, Spain) presented data on generating a co-stimulatory agonistic aptamer. This aptamer, which bound to both CD28 and multidrug resistant protein 1 (MRP-1), inhibited growth of tumours in mice (Soldevilla et al, 2016). Last in this session was Professor Victoria Calzada (Universidad de la República, Montevideo-Uruguay) who has linked technetium-99m and gallium-67 to an aptamer, sgc-8, through the chelators HYNIC and DOTA, and which showed good biodistribution and pharmacokinetics (Calzada et al, 2017).

**TOOLS/SELECTION/DESIGN (PART I)**

As technology is advancing rapidly, so too is the ability to develop aptamers more rapidly, or for advanced applications. As there are a number of exciting developments in this area, this topic was split across two sessions. The first part of this session, and the last for day 1 was chaired by Professor Marit Nilsen-Hamilton (Iowa State University, USA). Dr David Bunka (Aptamer Group Ltd, UK) started the session with a discussion on ‘Aptabind’ and their industrial application of protein purification. Their patented technology can be ‘tuned’ to the process to ensure that the process is not only efficient, but is also capable of purifying intact functional proteins. Next up was Dr Duncan Borthwick (Dynamic Biosensors GmbH, Germany) who has developed an electroswitchable biosurface to measure accurate binding kinetics of aptamers. Following this, Dr Sean Dembowski (University of Minnesota–Twin Cities, USA) discussed capillary electrophoresis as a means of enhancing the SELEX process for membrane proteins (Dembowski and Bowser 2017). Last of the session on day 1 was Mr John Goertz (University of Maryland, USA) who discussed a novel point-of-care device that uses a peroxidyme-based system allowing for assay automation (Goertz and White, 2017).

**TOOLS/SELECTION/DESIGN (PART II)**

Starting off day 2 and continuing the Tools/Selection/Design session, and chaired by Professor Fernando Pastor, was Dr Terry Steele (Nanyang Technological University, Singapore) who discussed the applications of multiple use fluorescent aptasensors, which could be used for rapid detection in real time field samples and effluents (Zhou et al, 2018). Dr Meltem Avci-Adali (University Hospital Tuebingen, Germany) discussed the use of aptamer functionalised hydrogels loaded with VEGF aptamers to capture endothelial cells to provide nutrients and oxygen delivery to cells (Guan et al, 2017). Last in this session was Professor Philip Johnson who discussed the complexities of two-site binding of the cocaine-binding aptamer, as well as the ATP-binding aptamer (Neves et al, 2017).
RIBOSWITCHES

The penultimate session focused on riboswitches and was chaired by Professor Ulrich Hahn. Riboswitches are RNA elements within an RNA transcript that modulate expression of proteins encoded by messenger RNA (Garst et al., 2011). Professor Beatrix Sueß discussed how aptamers can be applied as engineered riboswitches, providing several exciting recent examples from her research group (Berens et al., 2015). For example, the Sueß group has developed strategies to control exon skipping for splicing in cells, as well as tunable regulation of cell apoptosis. Dr Florian Groher (Technical University Darmstadt, Germany) then presented his work on the de novo selection and subsequent in vivo screening of a new synthetic riboswitch responsive to the antibiotic ciprofloxacin (Groher and Sueß, 2016; Groher et al., 2018). Both presentations emphasized the challenges in developing riboswitches from in vitro selected aptamers, and highlighted the need for improved in vivo screening systems following conventional SELEX. Dr Carlos Penedo (University of St Andrews, UK) presented elegant and insightful studies using single-molecule FRET for the structural analysis of the aptamer-expression platforms in riboswitch sequences (Blouin et al., 2015; Perez-Gonzalez et al., 2016). Completing the session on riboswitches, Professor Mario Mörl (Leipzig University, Germany), discussed some of the benefits and challenges of the modular composition of a standard riboswitch (Etzel and Mörl, 2017).

COMPUTATION

The final session, chaired by Dr David Bunka, focused on computational methods for analysing or selecting aptamers. This is a growing area of research, given the new applications available for in silico analysis of aptamers for both structure determination and selection. First, Dr Muslum Ilgu (Aptalogic Inc, USA) discussed his combined approach, using computational models and biochemical interrogation to determine aptamer 3D structure. Specifically, MC-Sym was used to predict a number of structures in silico. Next, 2-aminopurine (2-AP)-substituted aptamers were generated and analysed. With this modification, changes in fluorescence represented regions of structural flexibility and facilitated a rapid, and relatively inexpensive method for determining aptamer structure. Next, Dr Greg Penner (NeoVentures Biotechnology Inc, Canada) presented again, this time representing NeoVentures Biotechnology. At NeoVentures, they have developed an analysis of SELEX progression using next generation sequencing at each round of selection. Dr Penner spoke of SELEX as being a combination of math and chemistry. For example, as expected, as a selection experiment proceeds, the complexity of the pool decreases; however, NeoVentures tends to observe similar sequences beginning at rounds 7–10, and it is the trajectory throughout the selection experiments that is important when isolating/finding the best aptamer sequences. In the final talk of the conference, Dr Maureen McKeague (ETH Zürich, Switzerland) discussed her analysis of the past 23 years of SELEX experiments. Her analysis revealed that the best selection experiments (e.g., those with the highest affinity aptamers) were done at 37 degrees Celsius, with low magnesium concentrations, and using efficient methods for separating PCR double-stranded products (McKeague et al., 2015).

CONCLUSION

The conference concluded with Professor Dr Ulrich Hahn providing closing remarks prior to wishing everyone a safe journey home and a return for Aptamers 2018, which is the 5th Aptamers Symposium on 11th and 12th April 2018.

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COMPETING INTERESTS

None declared.

REFERENCES


