Advantages of a Dynamic Polygon for MHC Class I and II Immunopeptides

Francesco Pingitore¹, Michael Krawitzky¹, Josh Elias², Chris Adams¹; ¹Bruker Daltonics, San Jose, CA, USA, ²Chan Zuckerberg Biohub, Stanford - CA

ASMS 2021, FP 685

Introduction

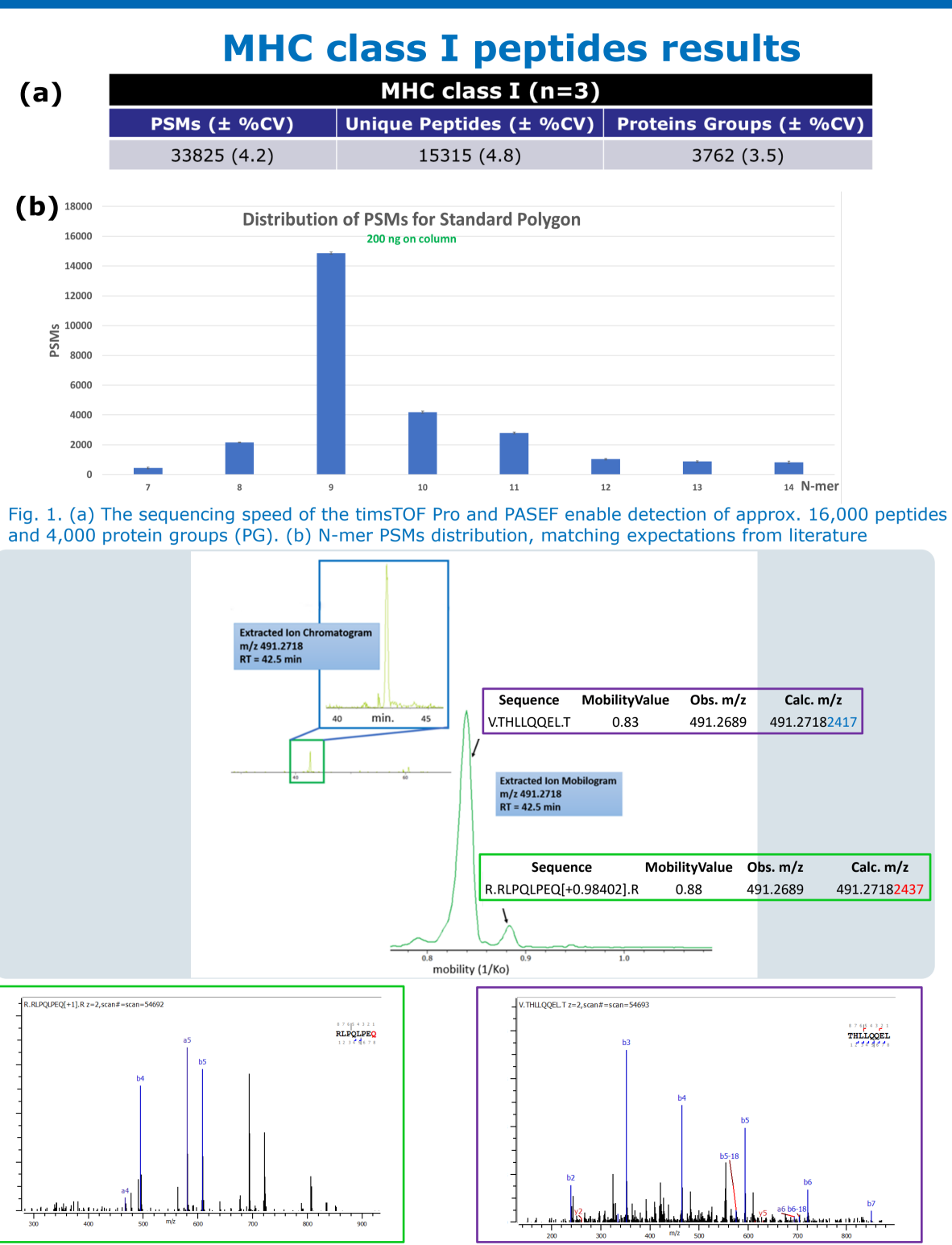
peptides modulate MHC-associated cell immunity and play a critical role in generating effective anti-tumor immune responses¹. Characterization of these peptides helps to generate therapeutic treatments and gain information on T cell mediated biomarkers. These peptides are challenging to characterize due to similar length, sequence conservation and lacking a defined termini when compared to peptides generated upon enzymatic digestion. To overcome these challenges, use of PASEF (Parallel Accumulation and Serial Fragmentation) enables to generate high quality peptide spectra and resolve coeluting and isobaric peptides. Moreover, the capability to easily tailor the mobility space enables preferential detection of groups and sub-groups of relevant peptides.

Methods

MHC class I peptides were separated on a 90 minutes gradient by nanoElute UPLC (Bruker Daltonics) on a 25 cm pulled emitter column (IonOpticks) and analyzed on a high resolution TIMS enabled QTOF instrument using the PASEF method (timsTOF Pro, Bruker Daltonics). An amount equal to 200 ng of estimated peptides were injected on column. For MHC class II peptides separation was performed on an EvoSep system equipped with an 8 cm performace column (a 60 samples per day method – 21 min gradient) and analyzed on the same instrument as previously described. An amount equal to 25 ng of estimated peptides was injected on the system. Data analysis was performed with PMI Byonic.

References

1) Lill et al.; Proteomics 2017, Volume 17, Issue 1-2: 1770010



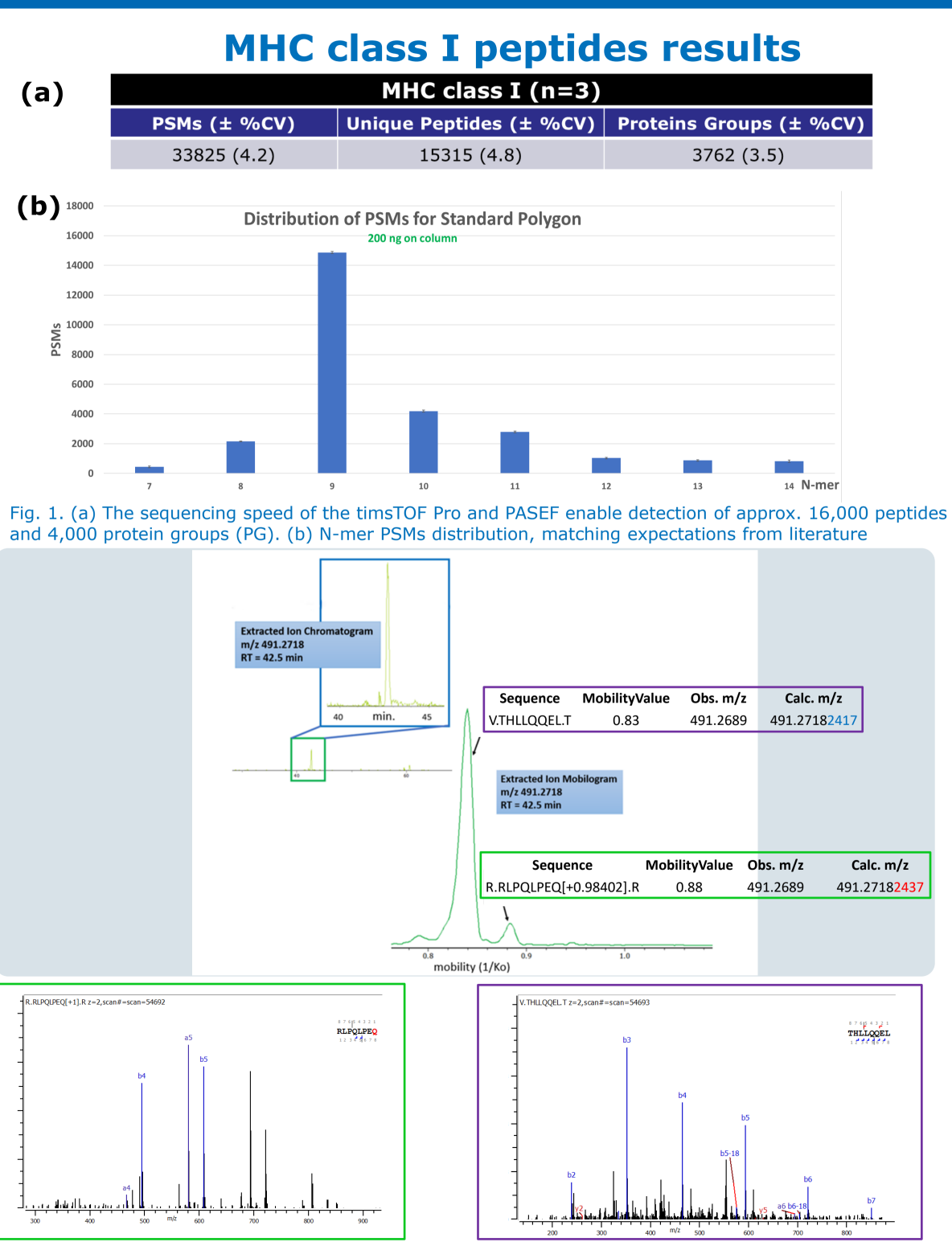
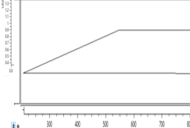
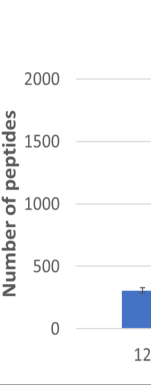


Fig. 2. Coeluting isobaric peptides at 42.56 min, are separated in the ion mobility dimension

60 SPD method – (amount injected on column 25 ng)							
	Polygon 200-1400 m/z		Polygon 250-950 m/z		Polygon 400-1300 m/z		
	Peptides	Proteins Groups	Peptides	Protein Groups	Peptides	Protein Groups	
Rep_1	1902	457	2196	514	1704	433	
Rep_2	1986	467	2559	558	1090	313	
Rep_3	1431	379	2280	528	1925	464	





N-mer	
12	
13	
14	
15	
16	
17	
18	

1317

1231

997

505

Fig. 4. N-mer distribution for middle polygon and mass range from figure 3 show N-mer varying from 14 to 16 are the most abundant species in the samples

1404

1279

955

541

1580

1419

1138

547

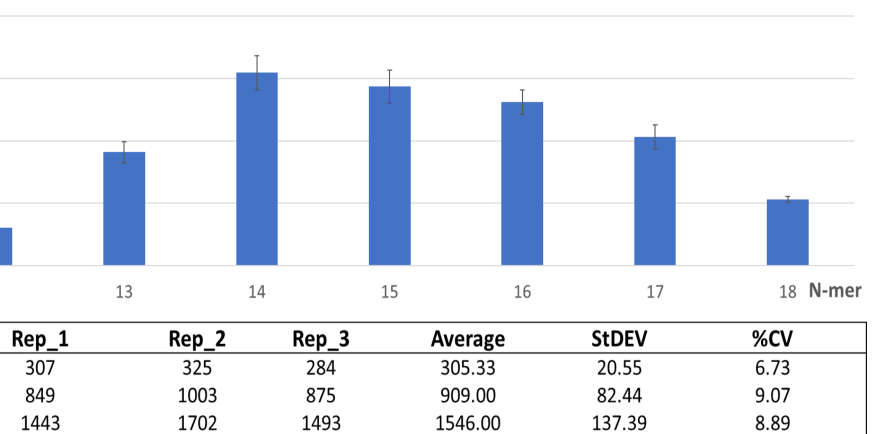
MHC class II peptides results

Fig. 3. Different polygon shapes and mass ranges used to maximize detection of peptides and PG; middle polygon generates best results

Analysis of MHC class I samples (figure 1) shows detection of a high number of peptides, with the unsurpassed sequencing speed and sensitivity of the timsTOFPro

Co-eluting peptides that are isobaric or have overlapping precursor ion isotope envelopes are resolved using Trapped Ion Mobility Spectrometry resulting in clean MS/MS spectra (figure 2)

Analysis of MHC class II samples with different settings (figure 3) shows dynamic parameters to maximize detection of peptides and PGs



1433.67

1309.67

1030.00

531.00

133.99

97.68

95.86

22.72

9.35

7.46

9.31

4.28

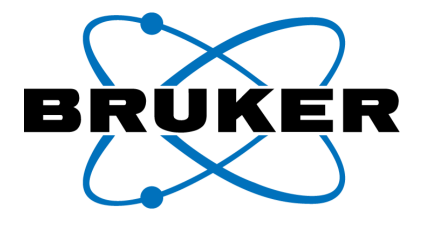
Graph and Table of major MHC II N-mer distribution

MHC class II peptides analyzed with EvoSep (Polygon 250 - 950)

More than 16000 peptides were confidently identified in a 90 min gradient, with high repeatability for MHC class I peptides.

timsTOF Pro





Results

Conclusions

PASEF on the timsTOF Pro with EvoSep separation identified more than 500 protein groups from a 21 min gradient separation, with just 25 ng on column (MHC class II peptides)

Extra dimension of separation provided by TIMS allowed resolution of co-eluting, isobaric peptides