Combinations of Autoantibodies Improve the Prediction of Timing of Onset of Future Rheumatoid Arthritis

- 1. University of Colorado Denver Anschutz Medical Campus.
- 2. Walter Reed National Military Medical Center, Bethesda, Maryland.
- 3. University of Nebraska, Omaha.

Introduction

Published data suggest that combinations of Anticitrullinated protein antibodies (ACPA) and Rheumatoid Factor (RF) are highly predictive of future rheumatoid arthritis (RA) as well as predictive of onset of RA within a relatively short time period. We have evaluated the role of combinations of ACPA and RF testing, and change over time, in predicting the time of onset of future clinically apparent RA.

Methodology

Using the Department of Defense Serum Repository we identified 215 RA cases. A mean of 3 pre-RA and 1 post-RA diagnosis serum samples were tested for RF immunoglobins (Ig) A, IgG, and IgM and anti CCP 2,3, and 3.1. The timing and trajectories of elevations of autoantibodies were evaluated. A gaptime cox regression model was used to develop hazard ratios for the risk of developing RA. Restricted meantime in state was also determined to predict time until RA diagnosis.

Results

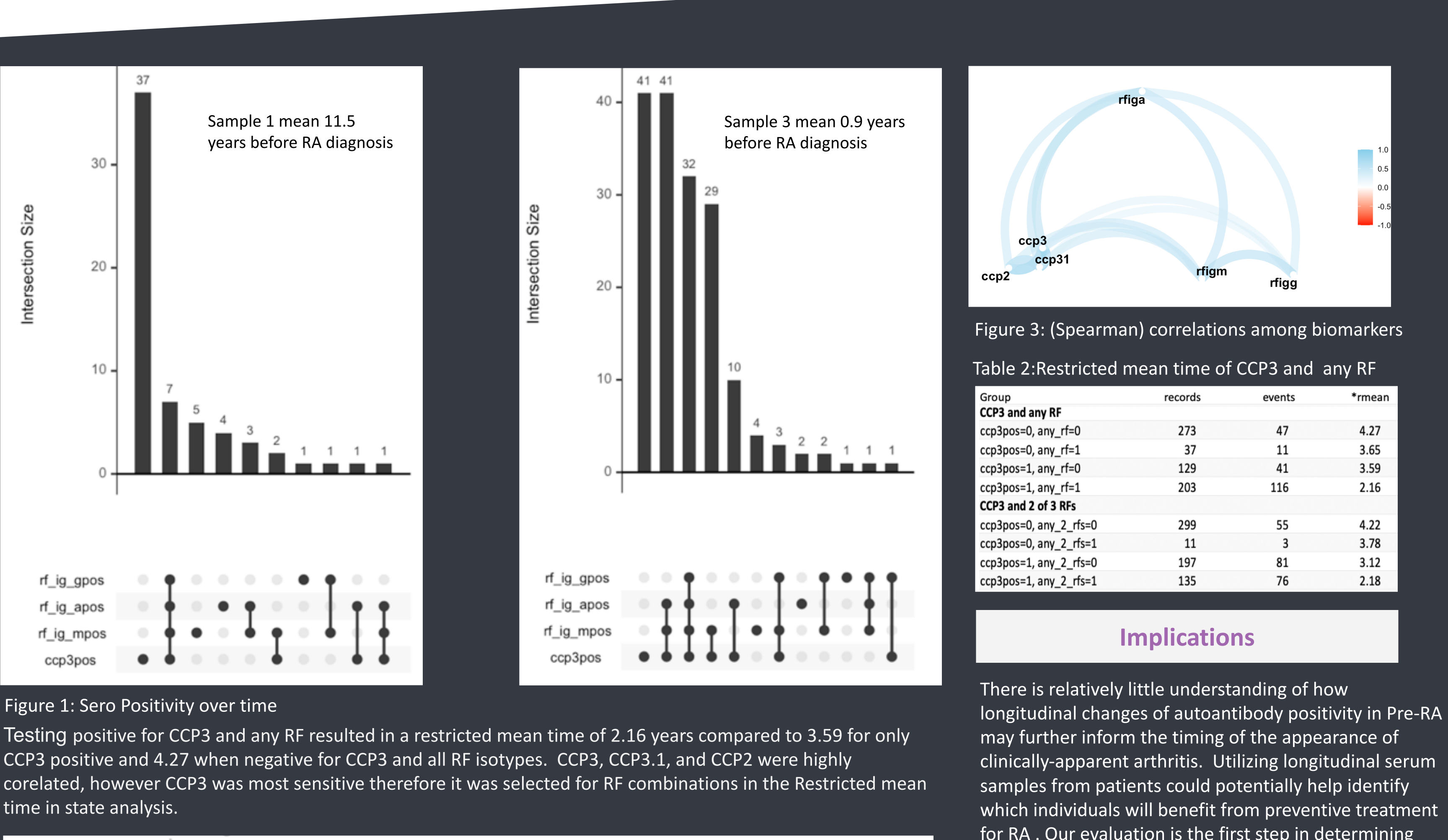
Table 1: Percent male and age at time of samples				
(N=215)				
age_earliest				
Mean (SD)	25.200 (6.305)			
Range	17.000 - 45.000			
age_latest				
Mean (SD)	38.014 (7.851)			
Range	20.000 - 58.000			
male	113 (52.6%)			

Controlling for age, gender, RFIgA and RFIgM status, if a subject had a positivity for either CCP2 or CCP3.1, they were at 3.3 times greater risk of developing RA compared to a subject who was not positive for either CCP2 or CCP3.1 (p < 0.001). A subject positive for RFIgA or RFIgM was at 1.6 times greater risk of developing RA (p = 0.002). These effects mean than subject testing positive for either CCP test and either RF test would be at 5.4 times greater risk than one who tested positive for neither.

Figure 1: Sero Positivity over time time in state analysis.

If a person has more positive sero markers it is more likely they will devolve RA, and the time until onset of clinically apparent RA symptoms will likely be shorter.

D.T. Bergstedt¹, R.A. Peterson¹, M.L. Feser¹, M C. Parish¹, J. D. Edison², T.R. Mikuls³, K.D. Deane¹



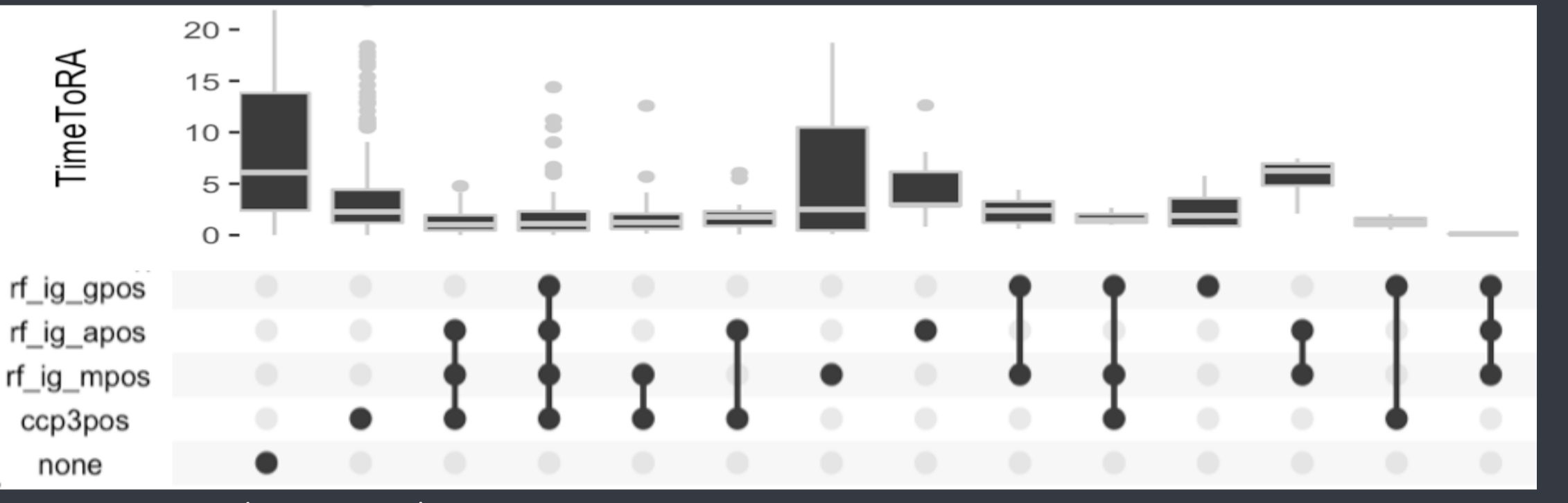


Figure 2: Restricted mean state by seropositivity

z.Resincted	mean time of	CCP5 and	ally Kr
	records	events	*rmear
l any RF			

	records	events	⁺rmean
l any RF			
:0, any_rf=0	273	47	4.27
:0, any_rf=1	37	11	3.65
:1, any_rf=0	129	41	3.59
:1, any_rf=1	203	116	2.16
2 of 3 RFs			
:0, any_2_rfs=0	299	55	4.22
:0, any_2_rfs=1	11	3	3.78
:1, any_2_rfs=0	197	81	3.12
:1, any_2_rfs=1	135	76	2.18

for RA. Our evaluation is the first step in determining the role of combinations of ACPA and RF testing, and change over time, in predicting the time of onset of future clinically apparent RA.

References

1.Rantapää-Dahlqvist S, et al. doi:10.1002/art.11223 2. Nielen MMJ, et al. doi:10.1002/art.20018 3. Deane KD, et al. doi:10.1002/art.27638 4. Van De Stadt LA, et al. doi:10.1136/annrheumdis-2012-202127 5. Rakieh C, et al. doi:10.1136/annrheumdis-2014-205227 6. Lingampalli N, et al. doi:10.1016/j.clim.2018.05.004 7. Deane K, et al.doi:10.1002/art.41417