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Development of forensic genomics research toolkits by the use of Massively Parallel Sequencing

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Epilogue

List of publications

Summary thesis

Nederlandse samenvatting

Dankwoord

Curriculum Vitae

List of publications

This thesis was based on the following publications:

- S Y Anvar, K J van der Gaag, J W F van der Heijden, M H A M Veltrop, R H A M Vossen, R H de Leeuw, C Breukel, H P J Buermans, J S Verbeek, P de Knijff, J T den Dunnen, J F J Laros. TSSV: a tool for characterization of complex allelic variants in pure and mixed genomes. *Bioinformatics* 02/2014
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