as median-high risk (table 1). The sensitivities and specificities are as follows for each tool: QRISK2 (19%, 93%), FRS (22%, 93%), mFRS (46%, 83%), QRISK3 (47%, 78%), SLECRE (61%, 63%), respectively. The tools were similar in negative predictive value, ranging from 89% (QRISK2) to 92% (SLECRE). The FRS and mFRS had the greatest c-statistics, both equalling 0.73, demonstrating the greatest predictive accuracy amongst the tools, while the QRISK3 and SLECRE had the lowest (0.67).

Conclusions While the mFRS performance was superior to the FRS, the QRISK3 did not outperform the mFRS. Although the SLECRE had the highest sensitivity, it had the lowest specificity, demonstrated by grouping the most cases and controls in the median-high risk category. Several factors are important to consider when deciding which risk assessment tools to utilize clinically: ease of use, sensitivity/specificity, and laboratory data accessibility. Thus, the mFRS continues to be a practical tool with a simple, intuitive scoring system appropriate for the ambulatory clinic setting based on the initial weighting of the FRS while adjusting for SLE.

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Cardiovascular Disease Risk Stratification by Risk Assessment Tool

Abstract 292 Figure 1

Background As the prototype of autoimmune disease, systemic lupus erythematosus (SLE) has complex and diverse clinical manifestations which may be harmful even life-threatening. It is pitiful that we can just passively respond to these serious complications. It will be a great advantage if the high-risk groups...
could be predicted and prevented with pre-treatment. The raising of risk prediction models depends on the collection of patient phenotypes, which are scattered in various forms and very cumbersome.

In this study, we collected the largest database of complete medical record of inpatients of lupus in China. The clinical phenotype database was generated by using natural language processing (NLP) techniques, then lupus nephritis (LN) prediction model was built.

**Methods** A total of 14,439 SLE patients were collected from the rheumatology and immunology departments of 13 Chinese tertiary hospitals in this study, including 13,062 females (90.46%), with an average age of 33.4 years, and the time span of EMR (Electronic Medical Records) was from October 28, 2001 to March 31, 2017. It includes basic information about patients, physical examination, inspection and diagnostic information, etc. We designed a hybrid NLP system combined NLP technical and expert knowledge at the same time, which was named as Deep Phenotyping System (DPS), to extract all the phenotypic information recorded in EMR. Based on these standard formatted entities, the machine learning and deep learning prediction methods are used to predict the LN in SLE.

**Results** The DPS efficiently processed EMR data, and its accuracy, precision, and recall were each greater than 93%. It extracted 73,794 entities from 14,439 SLE cases, each with time attributes, and produced 18,785,000,640 entities. Thus, a phenotype database was generated by using natural language processing (NLP) techniques, then lupus nephritis (LN) prediction model was built.

**Conclusions** The comprehensive SLE phenotype database constructed by NLP greatly improves the research efficiency of lupus clinical phenotype. We first proposed a predictive model of lupus nephritis, which is high applicability and efficiency. The experimental results of good close and open testing fully demonstrate the authenticity and practicality of this database. The research process and method based on real world data are also applicable to predict other important complications of lupus.

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**A NOVEL CD3/BCMA BISPECIFIC ANTIBODY SELECTIVELY KILLS PLASMA CELLS IN BONE MARROW OF HEALTHY INDIVIDUALS WITH IMPROVED SAFETY**

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**Background** Autoantibodies play an important role in the pathogenesis of systemic lupus erythematosus (SLE). Plasma cells secrete these autoantibodies but are unfortunately refractory to conventional immunosuppressive treatments. B-cell Maturation Antigen (BCMA) is exclusively expressed on mature B cells and, for that reason, is considered a good target for plasma cell depletion. Here, we analyze a promising therapeutic in development for Multiple Myeloma (MM), that could be used for the treatment for SLE. The antiCD3*BCMA bispecific antibody TNB-383B is in clinical trials for the treatment of MM. *In vitro* experiment showed that TNB-383B added to human bone marrow samples *ex vivo* induced a dose dependent lysis of BCMA-expressing plasma cells.

**Methods** Depletion of plasma cells by TNB-383B was tested using bone marrow samples extracted from bone retrieved of hip replacements. Human bone marrow mononuclear cells (n=10) were incubated with TNB-383B at increasing concentrations. The activity of TNB-383B was compared to a positive control antibody (PC), which has a BCMA-binding domain and an Fc region identical to TNB-383B, but binds to CD3 with stronger affinity. Bone marrow mononuclear cells were incubated for 18 hours with controls and TNB383B and samples were analyzed for plasma cell depletion (cells expressing CD19+, IgD-, IgM-, CD38++, CD27++) and T cell activation (CD69, CD107a, CD137, CD154). Cytokine production (IFN-, TNF, IL-2, IL-6, IL-12p70, IL-13, IL-4 and IL-10) was evaluated using multiplex.

**Results** TNB-383B effectively depleted plasma cells in bone marrow samples of healthy individuals undergoing hip replacement; more than 80% of BCMA expressing plasma cells were depleted after an overnight incubation with TNB-383B or a positive control. Analysis activation markers showed that TNB-383B activated T cells as evidenced by the expression of activation markers (CD69, CD154 or CD137) but only minimal cytokine production was observed.

**Conclusions** TNB-383B represents a novel immunotherapeutic with improved safety and efficacy for the treatment of pathogenic long-lived plasma cells. TNB-383B is in clinical trials for the treatment of Multiple Myeloma. TNB-383B could be an attractive treatment to prevent acute organ rejection in Panel Reactive Antibody (PRA) organ transplantation and autoimmune patients including Systemic Lupus Erythematosus. Teneobio anticipates to start testing TNB-383B in PRA patients by late 2019.

**Funding Source(s):** Teneobio, Inc.

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**IMMUNOPHENOTYPIC SUBGROUPS OF SLE DEFINED BY AUTOANTIBODIES, GENE EXPRESSION AND FLOW CYTOMETRIC ANALYSIS**

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**Background** SLE may be stratified according to a range of different immune assessments but the relationships between these are less well defined. MASTERPLANS is an MRC-funded consortium that seeks to identify immunophenotypic subgroups of patients that predict response to therapy. Our objective here was to analyse a clinically well-phenotyped patients using a suite of immune assessments and identify inter-relationships