Reduced False Positives in Autism Screening Via Digital Bio-markers Inferred from Deep Co-morbidity Patterns

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**Abstract**

Autism spectrum disorder (ASD) is a developmental disability associated with significant social and behavioral challenges. There is a need for tools that help identify children with ASD as early as possible1,2. Our current incomplete understanding of ASD pathogenesis, and the lack of reliable biomarkers hampers early detection, intervention, and developmental trajectories. In this study we develop and validate machine inferred digital biomarkers for autism using individual diagnostic codes already recorded during medical encounters. Our risk estimator identifies children at high risk with a corresponding area under the receiver operating characteristic curve (AUC) exceeding 80% from shortly after two years of age for either sex, and across two independent databases ofpatient records. Thus, we systematically leverage ASD co-morbidities - with no requirement of additional bloodwork, tests or procedures - to predict elevated risk during the earliest childhood years, when intervention is the most effective. Our methodology has superior performance to common questionnaires-based screenings such as the M-CHAT/F3, and has the potential to reduce socio-economic, ethnic and demographic biases. Further, by conditioning on the individual M-CHAT/F scores, we can either halve the false positives or boost sensitivity by over 50%, while maintaining specificity above 95%. Translated into practice, our algorithmic approach could significantly reduce the median diagnostic age for ASD, and also reduce long post-screen wait-times4 currently experienced by families for confirmatory diagnoses and access to evidence based interventions.

**INTRODUCTION**

Autism spectrum disorder is a developmental disability associated with significant social and behavioral challenges. Even though ASD may be diagnosed as early as the age of two5, children frequently remain undiagnosed until after the fourth birthday6. At this time, there are no laboratory tests for ASD, so a careful review of behavioral history, and a direct observation of symptoms is necessary7,8 for a clinical diagnosis. Starting with a positive initial screen, a confirmed diagnosis of ASD is a multi-step process that often takes 3 months to 1 year, delaying entry into time-critical intervention programs. While lengthy evaluations9, cost of care10, lack of providers11, and lack of comfort in diagnosing ASD by primary care providers11 are all responsible to varying degrees12, one obvious source of this delay is the number of false positives produced in the initial ASD-specific screening tools in use today. For example, the M-CHAT/F used commonly as a screening tool8,13, has an estimated sensitivity of 38.8%, specificity of 94.9% and Positive Predictive Value (PPV) of 14.6%3. Thus, currently out of every 100 children with ASD, M-CHAT/F flags about 39, and out of every 100 children it flags, about 85 are false positives, exacerbating wait times and queues12. Automated screening that might be administered with no specialized training, requires no behavioral observations, and is functionally independent of the tools employed in current practice, has the potential for immediate transformative impact on patient care.

While the neurobiological basis of autism remains poorly understood, a detailed assessment conducted by the US Centers for Disease Control and Prevention (CDC) demonstrated that children with ASD experience higher than expected rates of many diseases5. These include conditions related to dysregulation of immune pathways such as eczema, allergies, asthma, as well as ear and respiratory infections, gastrointestinal problems, developmental issues, severe headaches, migraines, and seizures14,15. In the present study, we exploit these co-morbidities to estimate the risk of childhood neuropsychiatric disorders on the autism spectrum. Using sequences of diagnostic codes from past doctor’s visits, our risk estimator reliably predicts an eventual clinical diagnosis — or the lack thereof — for individual patients. Thus, the key clinical contribution of this study is the formalization of subtle co-morbidity patterns as a reliable screening tool, and potentially improve wait-times for diagnostic evaluations by signiﬁcantly reducing the number of false positives encountered in initial screens in current practice.

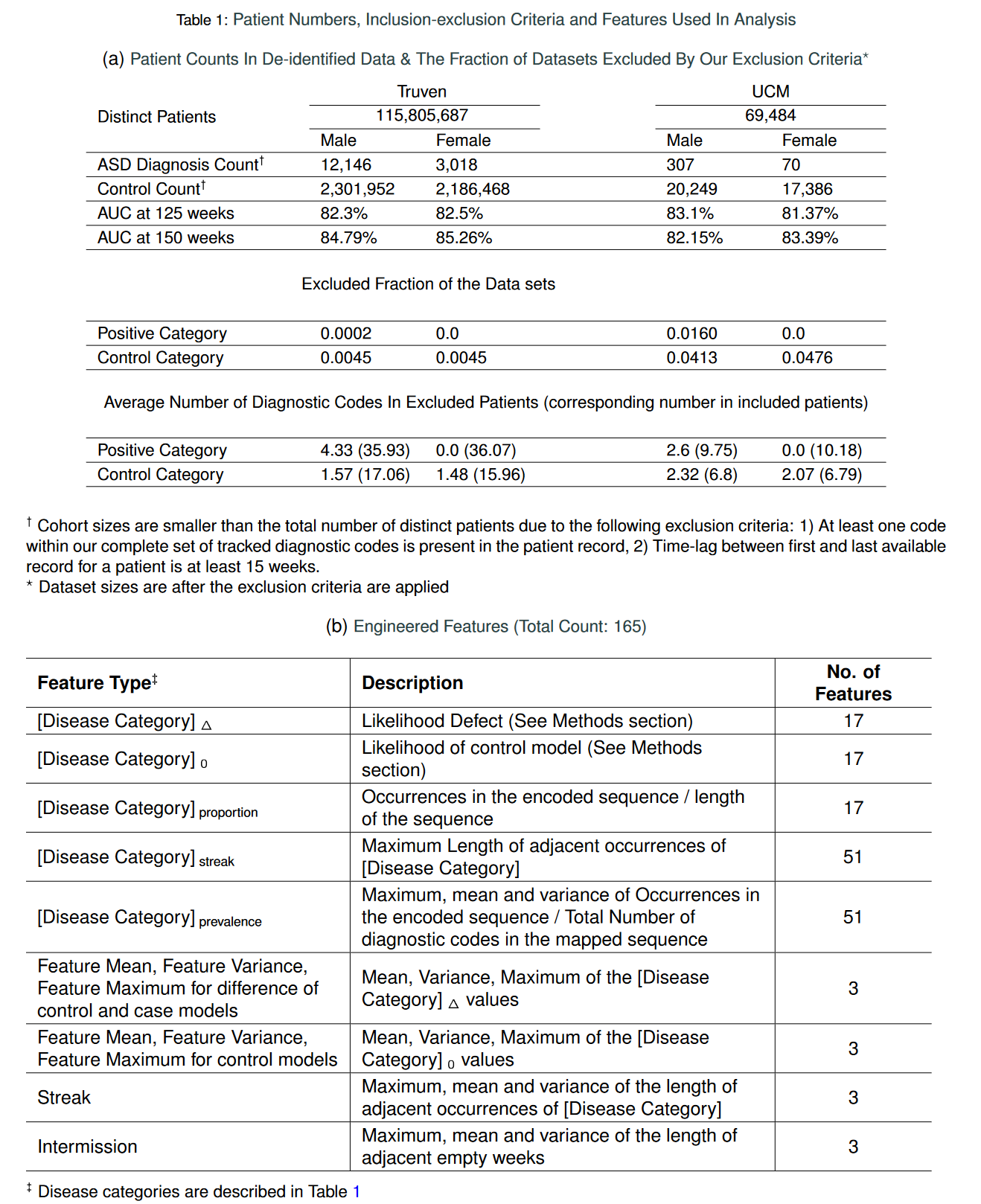
A screening tool that tracks the risk of an eventual ASD diagnosis, based on the information already being gathered during regular doctor’s visits, and which may be implemented as a fully automated background process requiring no time commitment from providers has the potential to reduce avoidable diagnostic delays at no additional burden of time, money and personnel resources. While still lacking the certainty of a diagnostic blood test, use of patterns emergent in the diagnostic history to estimate risk might help reduce the subjective component in questionnaire-based screening tools, resulting in 1) reduced effect of potential language and cultural barriers in diverse populations, and 2) possibly better identify children with milder symptoms8. Furthermore, being functionally independent of the M-CHAT/F, we show that there is a clear advantage to combining the outcomes of the two tools: we can take advantage of any population stratiﬁcation induced by the M-CHAT/F scores to signiﬁcantly boost combined screening performance (See Materials & Methods, and Supplementary text, section 8).

**MATERIALS & METHODS**

We view the task of predicting ASD diagnosis as a binary classiﬁcation problem: sequences of diagnostic codes are classiﬁed into positive and control categories, where “positive” refers to children eventually diagnosed with ASD, as indicated by the presence of a clinical diagnosis (ICD9 code 299.X) in their medical records. Of the two independent sources of clinical incidence data used in this study, the primary source used to train our predictive pipeline is the Truven Health Analytics MarketScan® Commercial Claims and Encounters Database for the years 2003 to 201218 (referred to as the Truven dataset). This US national database contains data contributed by over 150 insurance carriers and large self-insuring companies and is a culmination of over 4.6 billion inpatient and outpatient service claims and almost six billion diagnosis codes. We extracted histories of patients within the age of 0-9 years and excluded patients for whom: 1) At least one code of any available phenotypes is present, 2) Lag between ﬁrst and last available record for a patient should be at least 15 weeks. These exclusion criteria ensure that we are not considering patients who have too few observations to either train on. Additionally, during validation runs, we restricted the control set to patients observable in the databases to those whose last record is not before the ﬁrst 150 weeks of life. The characteristics of excluded patients are shown in Table 1a. We trained with over 30M diagnostic records (16,649,548 for males and 14,318,303 for females with 9,835 unique codes).

While the Truven database is used for both training and out-of-sample cross-validation with held-back data, our second independent dataset comprising de-identified diagnostic records for children treated at the University of Chicago Medical Center between the years of 2006 to 2018 (the UCM dataset) aids in further cross-validation. We considered children between the ages of 0 – 5 years, and applied the same exclusion criteria as the Truven dataset. The number of patients used from the two databases is shown in Table 1a. Our datasets are consistent with documented ASD prevalence and median diagnostic age (3 years in the claims database versus 3 years 10 months to 4 years in US17).

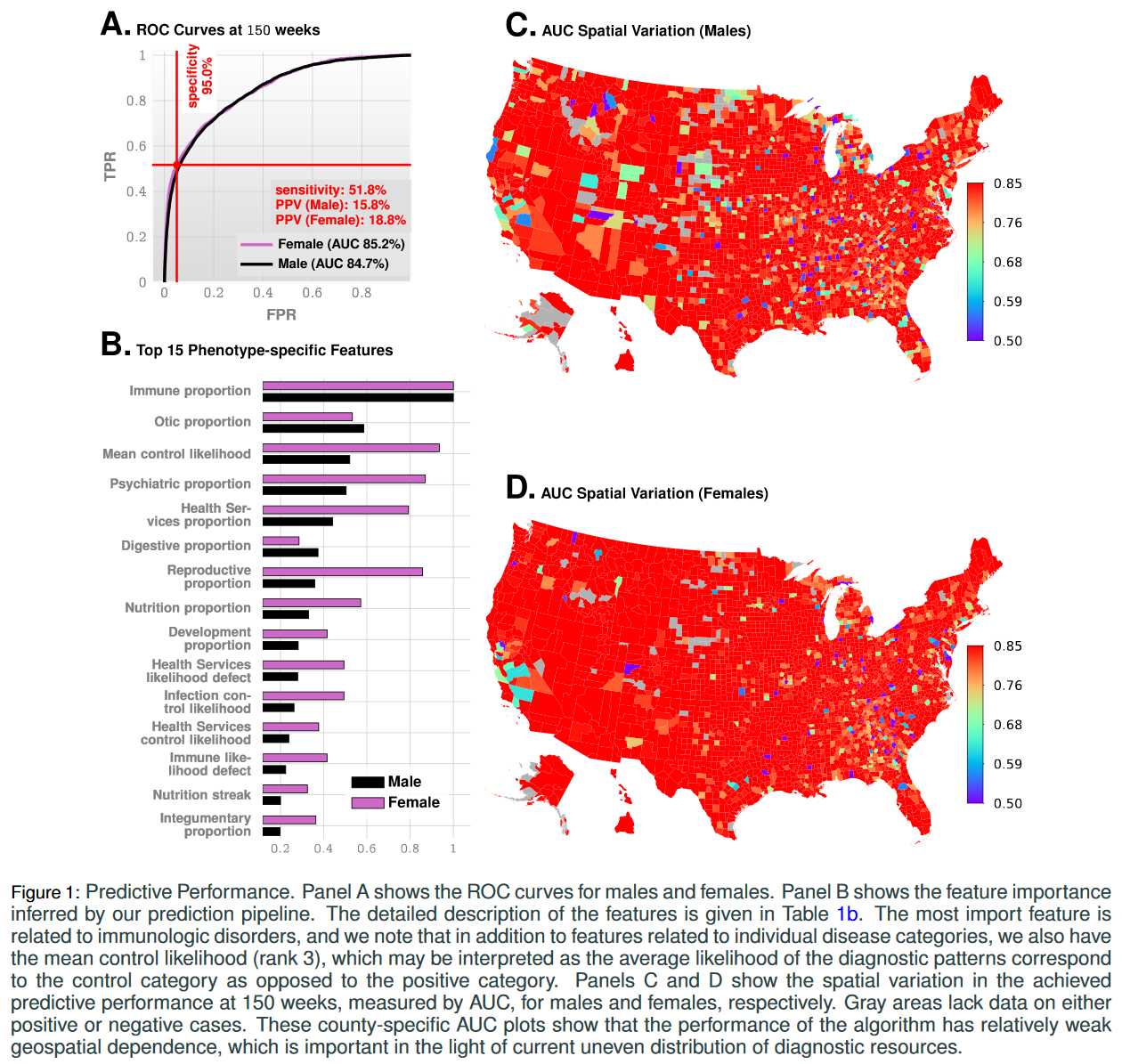
The signiﬁcant diversity of diagnostic codes (6,089 distinct ICD-9 codes and 11,522 distinct ICD-10 codes in total in the two datasets), along with the sparsity of codes per sequence and the need to make good predictions as early as possible, makes this a difﬁcult learning problem, where standard deep learning approaches do not sufﬁce (See Supp. Fig. 2). To address these issues, we proceed by partitioning the disease spectrum into 17 broad categories, e.g. infectious diseases, immunologic disorders, endocrinal disorders, etc. Each patient is then represented by 17 distinct time series, each tracking an individual disease category. At the population level, these disease-speciﬁc sparse stochastic time series are compressed into specialized Markov models (separately for the control

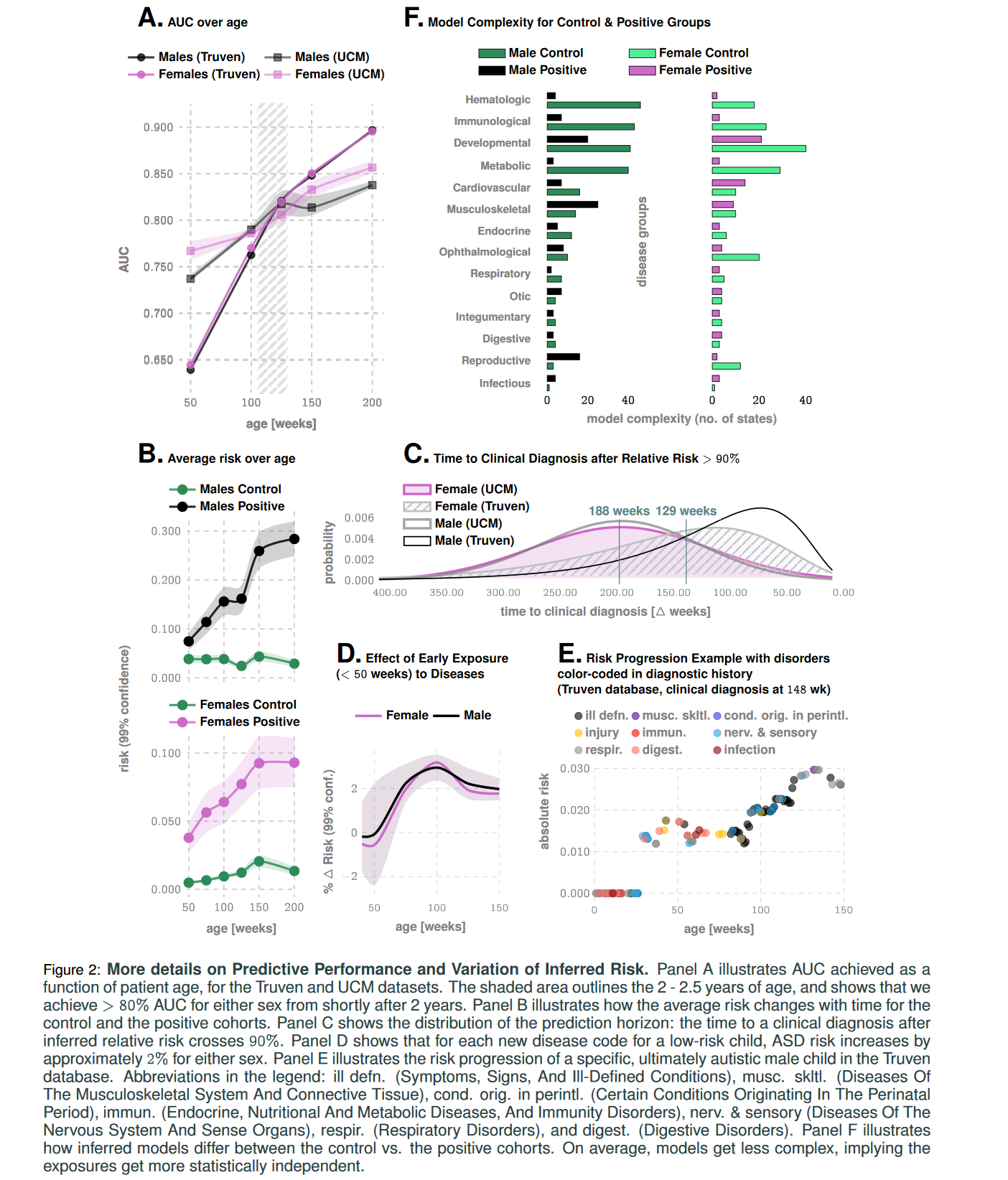
and the treatment cohorts) to identify the distinctive patterns pertaining to elevated ASD risk. As detailed in the Supplementary text, we use a novel approach to evaluate subtle deviations in stochastic observations known as the likelihood defect, to rigorously quantify similarity to the control vs the positive cohort for individual patients. In addition to the category specific Markov models, we use a range of engineered features that reflect various aspects of the diagnostic histories, including the proportion of weeks in which a diganostic code is generated, the maximum length of consecutive weeks with codes, the maximum length of weeks with no codes (See Tab. 1b for complete description), resulting in a total of 165 different features that are evaluated for each patient. With these inferred patterns included as features, we train a second level predictor that learns to map individual patients to the control or the positive groups based on their similarity to the identified Markov models of category-specific diagnostic histories, and the other engineered features (See Section 1 on detailed mathematical details in Supplementary text).

Our pipeline maps medical histories to a raw indicator of risk. Ultimately, to make crisp predictions, we must choose a decision threshold for this raw score. In this study, we base our analysis on maximizing the F1-score, defined as the harmonic mean of sensitivity and specificity, to make a balanced trade-off between Type 1 and Type 2 errors (See Supplementary text, Section 4). The relative risk is then defined as the ratio of the raw risk to the decision threshold, and a value >1 predicts a future ASD diagnosis. Our two step learning algorithm outperforms standard tools, and achieves stable performance across datasets strictly superior to documented M-CHAT/F.

The independence of our approach from questionnaire based screening implies that we can further boost our performance by conditioning choice of the sensitivity/specificity trade-offs on individual M-CHAT/F scores. In particular, we leverage the population stratification induced by M-CHAT/F to improve combined performance. We consider 4 sub-populations defined by M-CHAT/F score brackets3, and if the screen result is considered a positive(high risk, indicating the need for a full diagnostic evaluation) or a negative, i.e., low risk: 1) score 2 screening ASD negative, 2) score [3-7] screening ASD negative on follow-up, 3) score [3-7] and screening ASD positive on follow-up, and 4) score 8, screening ASD positive. (See SI-Table 3). The “follow-up” in the



context of M-CHAT/F refers to the re-evaluation of responses by qualified personnel. We use published data on the relative sizes and the prevalence statistics in these sub-populations3to compute the feasible conditional choices of our operating point to strictly supersede M-CHAT/F performance. Two limiting operating conditions are of special interest here, where we maximize PPV under some minimum specificity and sensitivity (denoted as the High Precision or the HP operating point), and where we maximize sensitivity under some minimum PPV and specificity (denoted as the High Recall or the HR operating point). Taking these minimum values of specificity, sensitivity, and PPV to be those reported for M-CHAT/F, we identify the set feasible set of conditional choices in a four-dimensional decision space that would outperform M-CHAT/F in universal screening. The results are shown in Fig. 3B.



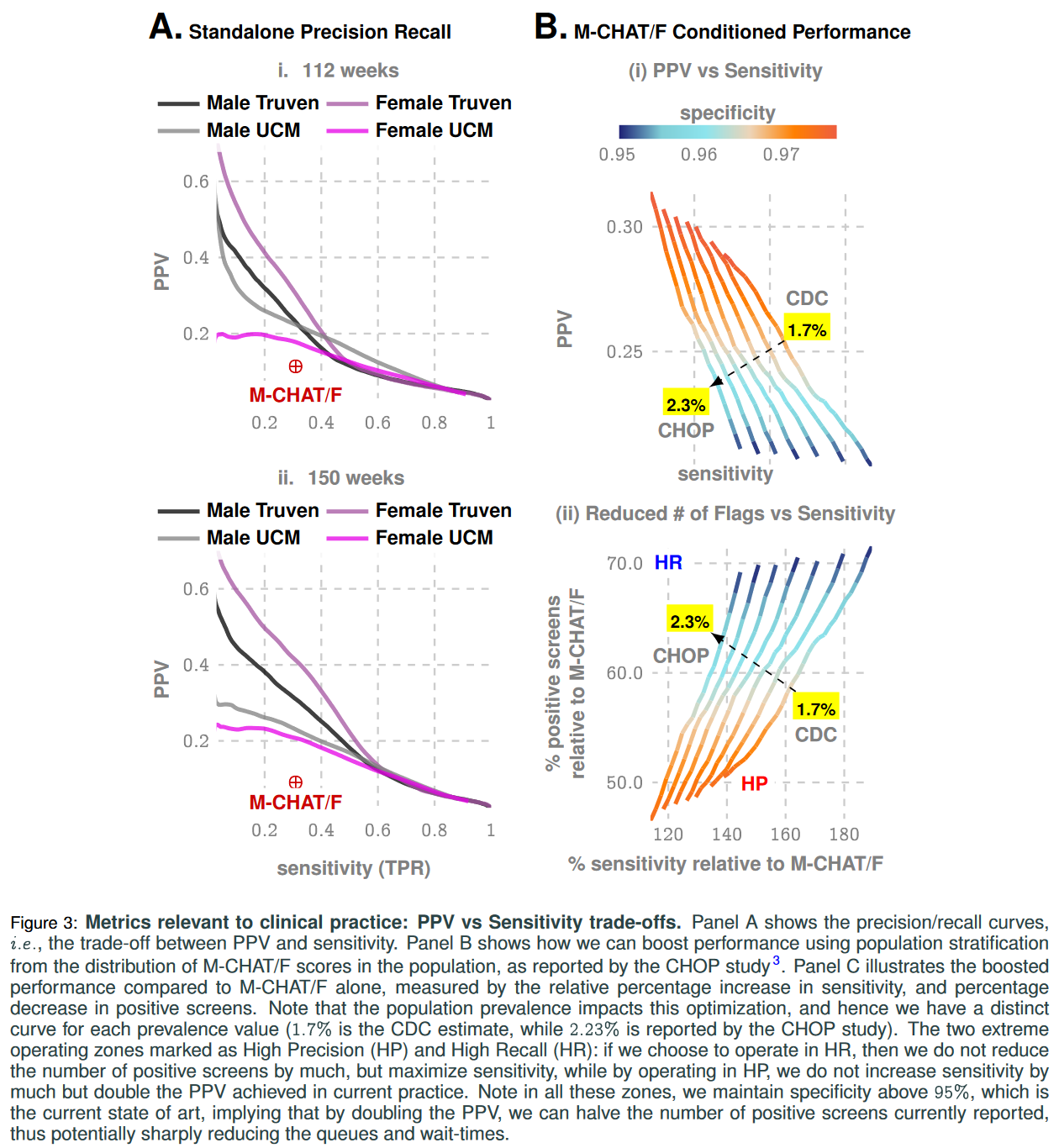
**RESULTS**

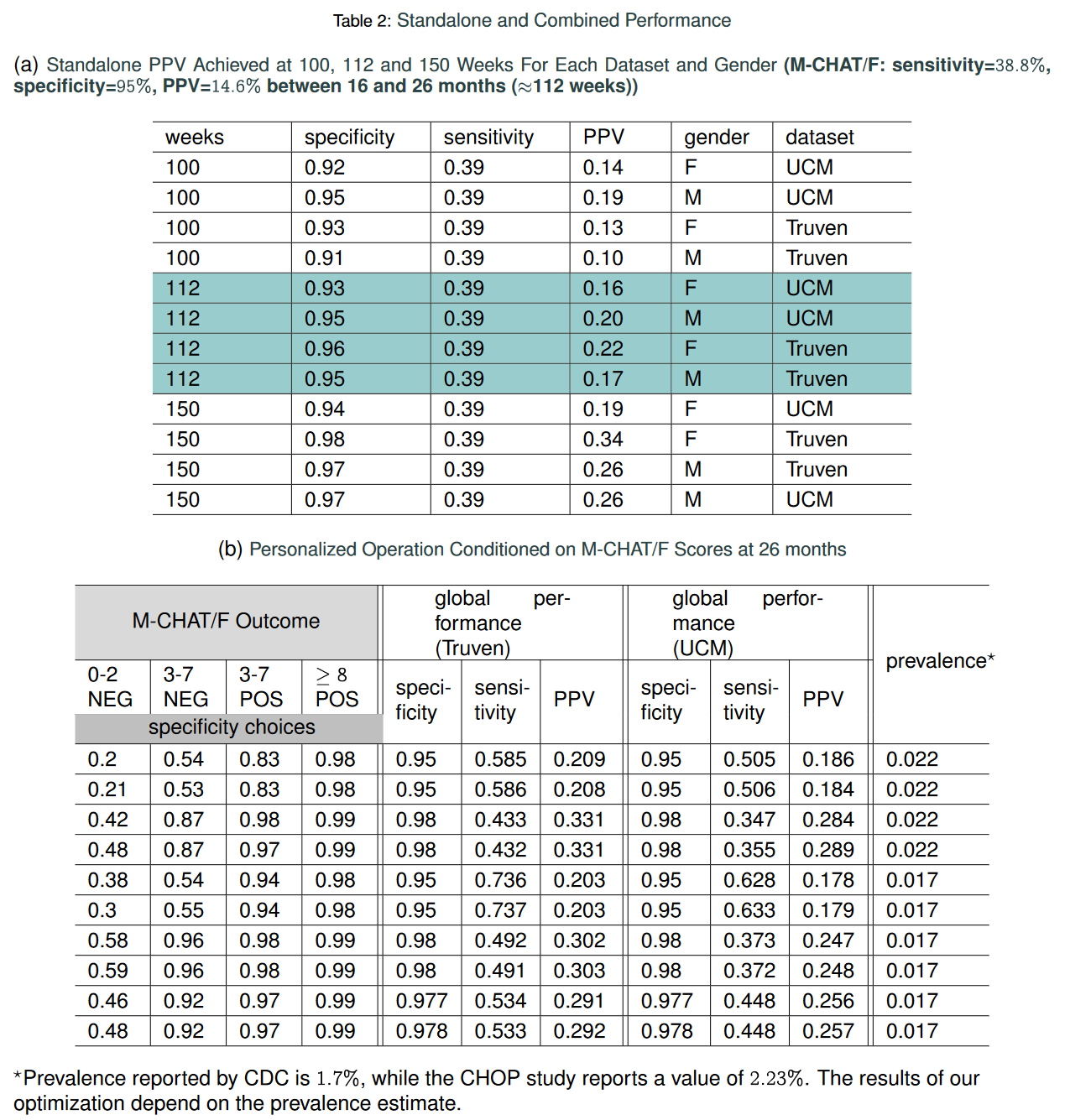
We measure our performance using several standard metrics including the AUC, sensitivity, specificity and the PPV. For the prediction of the eventual ASD status, we achieve an out-of-sample AUC of 82.3% and 82.5% for males and females respectively at 125 weeks for the Truven dataset. In the UCM dataset, our performance is comparable: 83.1% and 81.3% for males and females respectively (Fig. 1 and 2). Our AUC is shown to improve approximately linearly with patient age: Fig. 2A illustrates that the AUC reaches 90% in the Truven dataset at the age of four. Importantly, we train our pipeline on 50% of the Truven dataset, and use held back data from Truven, and the entirety of the UCM dataset for validation: No new training is done in the UCM dataset. Good performance on these independent datasets lends strong evidence for our claims. Furthermore, applicability in new datasets without local re-training makes it readily deployable in clinical settings.

Enumerating the top 15 predictive features (Fig. 1B), ranked according to their automatically inferred weights (the feature “importances”), we found that while infections and immunologic disorders are the most predictive, there is significant effect from all the 17 disease categories. Thus, the co-morbid indicators are distributed across the disease spectrum, and no single disorder is uniquely implicated (See also Fig. 2F). Importantly, predictability is relatively agnostic to the number of local cases across US counties (Fig. 1C-D) which is important in light of the current uneven distribution of diagnostic resources12,18 across states and regions.

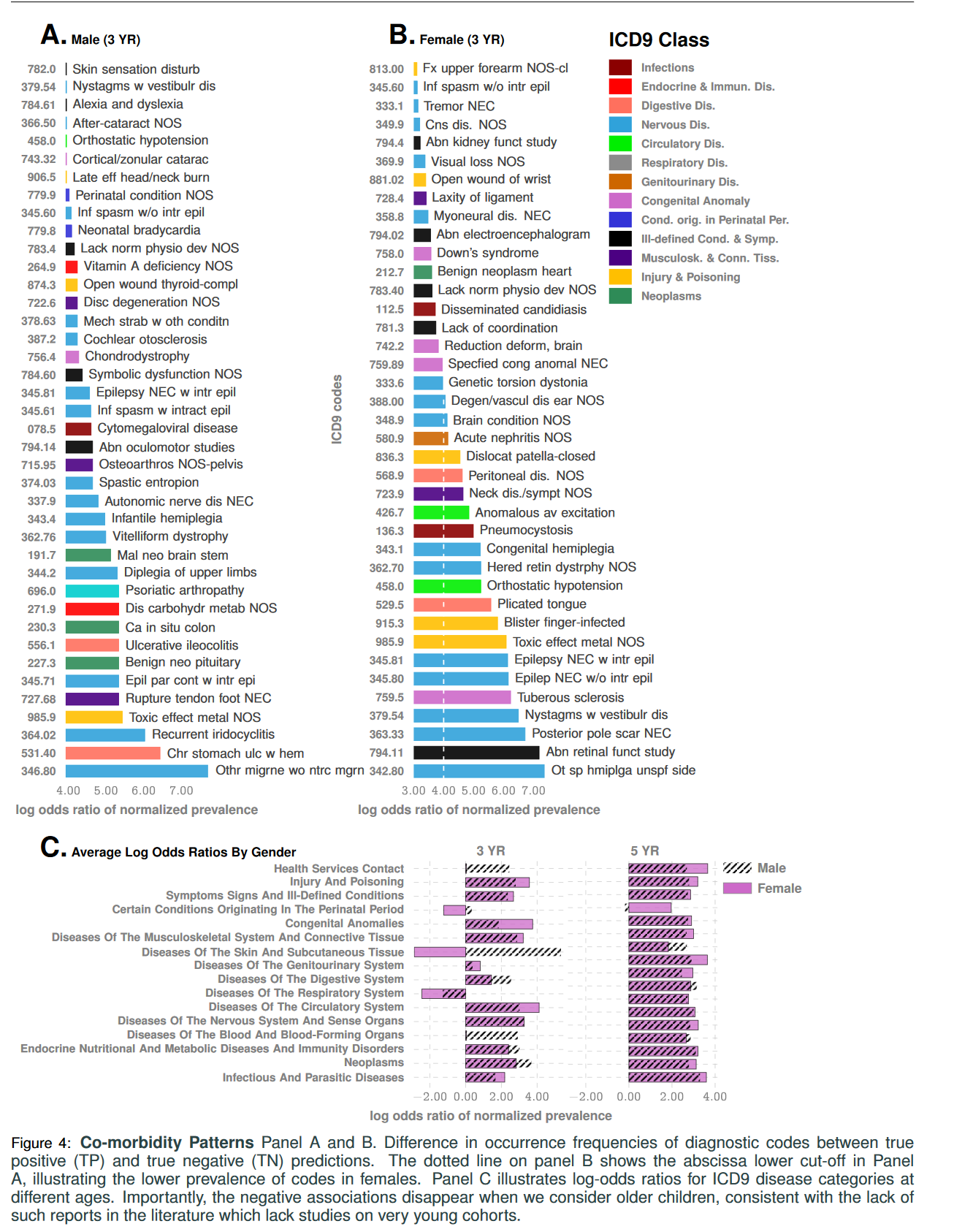
Unlike individual predictions which only become relevant over 2 years, the average risk over the populations is clearly different from around the first birthday (Fig. 2B), with the risk for the positive cohort rapidly rising. Also, we see a saturation of the risk after 3 years, which corresponds to the median diagnosis age in the database. Thus, if a child is not diagnosed up to that age, then the risk falls, since the probability of a diagnosis in the population starts to go down after this age. While average discrimination is not useful for individual patients, these reveal important clues as to



how the risk evolves over time. Additionally, while each new diagnostic code within the first year of life increases the risk burden by approximately 2% irrespective of sex (Fig. 2D), distinct categories modulate the risk differently, e.g., for a single random patient illustrated in Fig. 2F infections and immunological disorders dominate early, while diseases of the nervous system and sensory organs, as well as ill-defined symptoms, dominate the latter period.

Given these results, it is important to ask how much earlier can we trigger an intervention? On average, the first time the relative risk (risk divided by the decision threshold set to maximize F1-score, see Methods) crosses the 90% threshold precedes diagnosis by 188 weeks in the Truven dataset, and 129 weeks in the UCM dataset. This does not mean that we are leading a possible clinical diagnosis by over 2 years; a significant portion of this delay arises from families waiting in





queue for diagnostic evaluations. Nevertheless, since delays are rarely greater than one year12, we are still likely to produce valid red flags significantly earlier than the current practice.

Our approach produces a strictly superior PPV (exceeding M-CHAT/F PPV by at least 14% (14.1-33.6%) when sensitivity and specificity are held at comparable values (approx. 38% and 95%) around the age of 26 months (112 weeks). Fig. 3A and Table 2a show the out-of-sample PPV vs. sensitivity curves for the two databases, stratified by sex, computed at 100, 112 and 100 weeks. A single illustrative operating point is also shown on the ROC curve in Fig. 1C, where at 150 weeks, we have a sensitivity of 51.8% and a PPV of 15.8% and 18.8% for males and females respectively, both at a specificity of 95%.



Beyond standalone performance, independence from standardized questionnaires implies that we stand to gain substantially from combined operation. With the recently reported population stratification induced by M-CHAT/Fscores3 (SI-Table 4), we can compute a conditional choice of sensitivity for our tool, in each sub-population (M-CHAT/F score brackets: 0-2, 3-7 (negative assessment), 3-7 (positive assessment), and >8), leading to a significant performance boost. With such conditional operation, we get a PPV close to or exceeding 30% at the high precision (HP) operating point across datasets (>33% for Truven, >28% for UCM), or a sensitivity close toor exceeding 50% for the high recall (HR) operating point (>58% for Truven, >50% for UCM), when we restrict specificities to above 95% (See Table 2b, Fig. 3B(i), and SI-Fig. 7 in the supplementary text). Comparing with standalone M-CHAT/F performance (Fig. 3B(ii)), we show that for any prevalence between 1.7% and 2.23%, we can double the PPV without losing sensitivity at >98% specificity, or increase the sensitivity by ~50% without sacrificing PPV and keeping specificity = 94%.

**DISCUSSION**

In this study, we operationalize a documented aspect of ASD symptomology in that it has a wide range of co-morbidities14,15,19 occurring at above-average rates8. Association of ASD with epilepsy20, gastrointestinal disorders21–26, mental health disorders27, insomnia, decreased motor skills28, allergies including eczema21–26, immunologic19,29–35 and metabolic25,36,37 disorders are widely reported. These studies, along with support from large scale exome sequencing38,39, have linked the disorder to putative mechanisms of chronic neuroinflammation, implicating immune dysregulation and microglial activation31,34,40–43 during important brain developmental periods of myelination and synaptogenesis. However, these advances have not yet led to clinically relevant diagnostic biomarkers. Majority of the co-morbid conditions are common in the control population, and rate differentialsat the population level do not automatically yield individual risk44. Attempts at curating genetic biomarkers has also met with limited success. ASD genes exhibit extensive phenotypic variability, with identical variants associated with diverse individual outcomes not limited to ASD, including schizophrenia, intellectual disability, language impairment, epilepsy, neuropsychiatric disorders and, also typicaldevelopment45. Additionally, no single gene can be considered “causal” for more than 1% of cases of idiopathicautism46. In the absence of biomarkers, current screening in pediatric primary care visits uses standardized questionnaires to categorize behavior. This is susceptible to potential interpretative biases arising from language barriers, as well as social and cultural differences, often leading to systematic under-diagnosis in diverse populations8. In this study we use time-stamped sequence of past disorders to elicit crucial information on the developing risk of an eventual diagnosis, and formulate a screening protocol that is free from such biases, and yet significantly outperforms the tools in current practice. Going beyond screening performance, this approach provides a new tool to uncover clues to ASD pathobiology. Perhaps this vulnerability to diverse immunological, endocrinological and neurological impairments reflects how allostatic loads of medical stress get under the skin and disrupt key regulators of CNS organization and synaptogenesis. Charting individual disorders in the co-morbidity burden further reveals novel associations in normalized prevalence — the odds of experiencing a specific disorder, particularly in the early years (age <3 years), normalized over all unique disorders experienced in the specified time-frame. We focus on the true positives in the positive cohort and the true negatives in the control cohort to investigate patterns that correctly disambiguate ASD status. On these lines Fig. 4 and SI-Fig. 4 in the supplementary text outline two key observations: 1) negative associations: some diseases that are negatively associated with ASD with respect to normalized prevalence, i.e., having those codes relatively over-represented in one’s diagnostic history favors ending up in the control cohort, 2) impact of sex: there are sex-specific differences in the impact of specific disorders, and given a fixed level of impact, the number of codes that drive the outcomes is significantly more in males (Fig. 4A vs B).Some of the disorders that show up in Fig. 4, panels A and B are surprising, e.g., congenital hemiplegia or diplegia of the upper limbs indicative of either cerebral palsy (CP) or a spinal cord/brain injury, neither of which has a direct link to autism. Since only about 7% of the children with cerebral palsy (CP) are estimated to have a co-occurring ASD47,48, and with the prevalence of CP significantly lower (1 in 352 vs 1 in 59 for autism), it follows that only a small number of children (approximately 1.17%) with autism have co-occurring CP. Thus, with significantly higher prevalence in children diagnosed with autism compared to the general population (1.7% vs 0.28%), CP codes show up with higher odds in the true positive set. Also, SI-Fig. 4A shows that the immunological, metabolic, and endocrine disorders are almost completely risk-increasing. In contrast, respiratory diseases (panel B) are largely risk-decreasing. On the other hand, infectious diseases have roughly equal representations in the risk-increasing and risk-decreasing classes (panel C). The risk-decreasing infectious diseases tend to be due to viral or fungal organisms, which might point to the use of antibiotics in bacterial infections, and the consequent dysbiosis of the gut microbiota23,37 as a risk factor. Any predictive analysis of ASD must address if we can discriminate ASD from general developmental and behavioral disorders. The DSM-5 established a single category of ASD to replace the subtypes of autistic disorder, Asperger syndrome, and pervasive developmental disorders8. This aligns with our use of diagnostic codes from ICD9 299.X as specification of an ASD diagnosis, and use standardized mapping to 299.X from ICD10 codes when we encounter them. For other psychiatric disorders, we get high discrimination reaching AUCs over 90% at 100 – 125 weeks of age (SI-Fig. 3A), which establishes that our pipeline is indeed largely specific to ASD. We carried out a battery of tests to ensure that our results are not significantly impacted by class imbalance (since our control cohort is orders of magnitude larger) or systematic coding errors (See Methods), e.g., we verified that restricting the positive cohort to children with at least two distinct ASD diagnostic codes in their medical histories instead of one, has little impact on out-of-sample predictive performance (SI-Fig. 3B). Can our performance be matched by simply asking how often a child is sick? We found that the density of codes in a child’s medical history is indeed somewhat predictive of a future ASD diagnosis, with the AUC 75% in the Truven database at 150 weeks (See SI-Fig. 3, panel D in the supplementary text). This is expected, since children with autism do indeed have higher rates of co-morbidities. However, it does not have stable performance across databases, and has no significant effect once the rest of the features are combined. As a key limitation to our approach, automated pattern recognition might not reveal true causal precursors. The relatively uncurated nature of the data does not correct for coding mistakes by the clinician and other artifacts, e.g. a bias towards over-diagnosis of children on the borderline of the diagnostic criteria due to clinicians’ desire to help families access service, and biases arising from changes in diagnostic practices over time49. Discontinuities in patient medical histories from change in provider-networks can also introduce uncertainties in risk estimates, and socio-economic status of patients which impact access to healthcare might skew patterns in EHR databases. Despite these limitations, the design of a questionnaire-free component to ASD screening that systematically leverages co-morbidities has far-reaching consequences, by potentially slashing the false positives and wait-times, as well as removing systemic under-diagnosis issues amongst females and minorities. Future efforts will attempt to realize our approach within a clinical setting. We will also explore the impact of maternal medical history, and the use of calculated risk to trigger blood-work to look for expected transcriptomic signatures of ASD. Finally, the analysis developed here applies to phenotypes beyond ASD, thus opening the door to the possibility of general comorbidity-aware risk predictions from electronic health record databases.



**Data Sharing**

Software implementation of the pipeline is available at: <https://github.com/zeroknowledgediscovery/ehrzero>, and installation in standard python environments may be done from <https://pypi.org/project/ehrzero/>.

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