TITLE PAGE

Title: A normative chart for cognitive development in a genetically selected population.

Authors: A.M. Fiksinski¹,²,³, C.E. Bearden⁴, A.S. Bassett²,³,⁵,⁶, R.S. Kahn⁷¹, J.R. Zinkstok¹, S. R. Hooper⁸, W. Tempelaar², the 22q11DS International Consortium on Brain and Behavior#, J.A.S. Vorstman*¹,²,³, E.J. Breetvelt*⁹

# Group authorship: complete list of contributing authors to be listed in Supplemental Materials 2.
* These authors contributed equally to this work.

Author affiliations:
¹ Department of Psychiatry, Brain Center, University Medical Center Utrecht, Utrecht, The Netherlands.
² Centre for Addiction and Mental Health, Toronto, Ontario, Canada.
³ The Dalglish Family 22q Clinic for 22q11.2 Deletion Syndrome, Toronto General Hospital, University Health Network, Toronto, Ontario, Canada.
⁴ Departments of Psychiatry and Biobehavioral Sciences and Psychology, Semel Institute for Neuroscience and Human Behavior, UCLA, Los Angeles, CA
⁵ Division of Clinical and Metabolic Genetics, The Hospital for Sick Children, Toronto, Ontario, Canada & Medical Genetics and Genomics Residency Training Program, University of Toronto, Toronto, Ontario, Canada
⁶ Toronto General Research Institute and Campbell Family Mental Health Research Institute, Toronto, Ontario, Canada.
⁷ Department of Psychiatry, Icahn School of Medicine at Mount Sinai, New York, NY, USA.
⁸ Department of Allied Health Sciences, School of Medicine, University of North Carolina-Chapel Hill, Chapel Hill, North Carolina, USA.
⁹ Department of Psychiatry, University of Toronto, Toronto, Ontario, Canada.
¹⁰ Program in Genetics and Genome Biology, Research Institute, and Department of Psychiatry, The Hospital for Sick Children, Toronto, Ontario, Canada.

Corresponding author:
Ania M. Fiksinski, MSc.
University Medical Center Utrecht, KA. B01107
Heidelbergaan 100
P.O. Box 85500
3508 GA, Utrecht
The Netherlands
T +31 88 7553275
E a.m.fiksinski@umcutrecht.nl

Word count: 3828

Abstract word count: 275
Certain pathogenic genetic variants impact neurodevelopment and cause deviations from typical cognitive trajectories. Understanding variant-specific cognitive trajectories is clinically important for informed monitoring and identifying patients at risk for comorbid conditions. Here, we demonstrate a variant-specific normative chart for cognitive development for individuals with 22q11.2 deletion syndrome (22q11DS). We used IQ data from 1365 individuals with 22q11DS to construct variant-specific normative charts for cognitive development (Full Scale, Verbal, and Performance IQ). This allowed us to calculate Z-scores for each IQ datapoint. Then, we calculated the change between first and last available IQ assessments (delta Z-IQ-scores) for each individual with longitudinal IQ data (n = 708). We subsequently investigated whether using the variant-specific IQ-Z-scores would decrease required sample size to detect an effect with schizophrenia risk, as compared to standard IQ-scores. The mean Z-IQ-scores for FSIQ, VIQ, and PIQ were close to 0, indicating that participants had IQ-scores as predicted by the normative chart. The mean delta-Z-IQ-scores were equally close to 0, demonstrating a good fit of the normative chart and indicating that, as a group, individuals with 22q11DS show a decline in IQ-scores as they grow into adulthood. Using variant-specific IQ-Z-scores resulted in 30% decrease of required sample size, as compared to the standard IQ-based approach, to detect the association between IQ-decline and schizophrenia (p<0.01). Our findings suggest that using variant-specific normative IQ data significantly reduces required sample size in a research context, and may facilitate a more clinically informative interpretation of IQ data. This approach allows identification of individuals that deviate from their expected, variant-specific,
trajectory. This group may be at increased risk for comorbid conditions, such as schizophrenia in the case of 22q11DS.

**Key words:** Cognitive development, IQ, high-risk, pathogenic genetic variant, 22q11DS, schizophrenia, normative chart
1. INTRODUCTION

Over the past two decades, a growing list of genetic variants associated with clinical phenotypic outcomes has emerged, including cognitive trajectories that deviate from what is typical in the general population[1-3]. In the general population, the age-adjusted level of cognitive functioning is generally stable over the lifespan; i.e., the IQ curve, where obtained scores are age-adjusted, is expected to be a virtually constant line over the years [4]. A divergent trajectory may be part of the developmental impact of an underlying pathogenic genetic variant. Examples include early cognitive decline and loss of acquired skills in the case of Rett’s syndrome [5,6], or early onset dementia in the case of Down’s syndrome [7-10]. General cognitive functioning is the term we use in this article to reflect the important human quantitative trait that accounts for much of the variation in diverse cognitive abilities, including intellectual functioning, and can be operationalized as the commonly used Intelligence Quotient (IQ) [11-13].

Populations of carriers of pathogenic variants that impact neurodevelopment would benefit from a better understanding of variant-specific cognitive trajectories. To that end, ideally variant-specific (age-) normative reference data are obtained, allowing for the comparison of an individual’s performance to the group’s indices over time and potentially helpful in setting realistic expectations regarding (future) performance. This is analogous to the significantly improved accuracy and clinical relevance of monitoring physical growth in an individual with Down’s syndrome when using normative physical growth data from studies of individuals with Down’s syndrome [14,15]. When using norm data obtained from the general population, a child with Down’s syndrome may be
considered growth-delayed, whereas in reality their growth trajectory may be as expected for someone with this genetic condition.

In a similar way, genetic subgroup-specific normative data on cognitive development may be highly informative. Such cognitive norm charts may be relevant for both research and clinical purposes as they allow the identification of individuals who deviate from what is a typical trajectory given the genetic variant and potentially, monitoring effects of interventions over time. For example, when an individual does not follow their expected IQ trajectory; i.e., deviates from their IQ curve, this may indicate underlying brain-related pathology, warranting additional examinations. A parallel may be drawn to how in a child who deviates from their expected physical growth curve a diagnostic work-up is warranted that could help identify the cause (e.g., endocrine problems), and potentially inform treatment strategies (e.g., growth hormones) [16].

The 22q11.2 deletion syndrome (22q11DS) is a genetic condition associated with aberrant neurodevelopmental outcomes [17]. It is the most common chromosomal microdeletion disorder, estimated to result from (in ~85% of cases de novo) non-homologous meiotic recombination events occurring in approximately 1 in every 1,000 fetuses [18]. 22q11DS has a highly variable phenotypic expression [19-22], including various levels of cognitive functioning with differing developmental trajectories that, on average, appear to display a mild downward trend [17,23]. Individuals with 22q11DS also have a 25-fold increased risk for developing schizophrenia, making it the strongest single molecular genetic risk variant for psychotic disorders [24]. We have previously reported that the subgroup of individuals with a cognitive decline steeper than average in this population had an even further elevated risk for schizophrenia [25]. Here, we
aim to generate a 22q11DS-specific normative chart for IQ to be used as a reference in both clinical and research settings. We will demonstrate how a normative chart for cognitive development in a genetically defined population can be reliably established and provide potential directions for its future utility.

2. METHODS

2.1 Participants and instruments

Data on 1789 individuals with a confirmed 22q11.2 microdeletion were collected from 22 different sites as part of the international Brain and Behavior Consortium on 22q11DS [26] [27]. For this study, we included individuals who had at least one Wechsler IQ assessment available and were between the ages of 6 and 38 years, resulting in a total number of participants of 1365 (76.3%). Of those, 657 individuals (48.1%) had one assessment available, and we refer to this subgroup as the baseline sample. 708 (51.9%) individuals had two or more IQ assessments available, and we refer to this subgroup as the longitudinal sample (see also Figure 1). All individuals, and when appropriate their legal guardians, provided informed consent and the study was approved by the local institutional research ethics boards of each site.

Level of overall intellectual functioning (IQ) was assessed using age-appropriate Wechsler scales (see also Table 1) [28-34] and all IQ-data underwent extensive quality control. Clinical diagnoses of schizophrenia spectrum disorders were made by experienced clinicians in accordance with the Diagnostic and Statistical Manual of Mental Disorders, Fourth [35] or Fifth edition [36]. Positive psychotic symptoms were assessed using standardized clinical interviews, including the Positive and Negative Syndrome Scale (PANNS [37]), the Comprehensive Assessment of At Risk Mental States
(CAARMS [38]), the Structured Interview for Psychotic Syndromes (SIPS [39]), and the Schedule for Affective Disorders and Schizophrenia for School-Age Children – Present and Lifetime Version (K-SADS-PL [40]).

2.2 Data analysis

The data analysis for this study consisted of two steps. First, we constructed the normative charts for IQ and second, we used the available longitudinal data to calculate the difference (delta) scores between the first and last available IQ assessments (see also Figure 1). All data quality control and statistical analyses were conducted in R 3.6.2 GUI 1.70.

2.2.i. Normative chart

To construct the normative chart for IQ, we used all available IQ datapoints (n = 2512) from all participants with at least 1 IQ-assessment available (n = 1365; all 22q11DS individuals from the IBBC). We used polynomial regression models of the 1st, 2nd, 3rd and 4th order and we used the Akaike and the Bayesian Information Criterion (AIC and BIC) to determine the best fit. Furthermore, we checked basic assumptions for polynomial regressions, including multivariate normality and homoscedasticity, by examining the distribution of the residuals and the residual variance of the final model.

Subsequently, we used the coefficients derived from the best fit to determine the normative IQ chart. This normative chart enabled us to calculate a (standardized) Z-score for each individual IQ-point, and thereby identify how much individuals deviated from the average IQ in this population at a certain timepoint, given their age. We applied the same strategy for all basic summary scores: Full Scale IQ (FSIQ), Verbal IQ (VIQ), and Performance IQ (PIQ).
2.2.ii Delta Z-scores

For those individuals with more than one IQ point available (n = 708, 51.9%), we determined delta-Z-scores by calculating the difference between the Z-score corresponding to the first available IQ measurement and that of the last IQ-measurement \( (\text{Last IQ Z-score} - \text{first IQ Z-score} = \text{delta-Z-score}) \). The average delta-Z score across all participants provides an indication of the extent to which individuals follow, on average, their expected trajectory as predicted by the normative chart. In addition, we examined the distribution of the delta-Z-scores.

2.2.iii Post-hoc

Post-hoc, we investigated whether using the delta-Z-score, as compared to the standard (population-normed) IQ-scores would result in a decrease of required sample size to detect the previously reported association between IQ-decline and schizophrenia risk [25]. To this end, we used odds ratios (ORs) from two regression models: both models had schizophrenia status (yes/no) as the dependent variable and baseline VIQ (we chose to focus on VIQ and VIQ-decline as this component of IQ had the strongest association with schizophrenia risk [25]), sex, age, and time-interval as covariates. In the (variant-specific) Z-based model the main (binary) independent variable was VIQ-Z-decline (yes/no; based on a cut-off of -0.5 SD in delta Z-score). In the parallel model the main independent variable was VIQ-decline (yes/no; based on a cut-off of -7.5 IQ-points (i.e., -0.5 SD) in absolute (population normed) VIQ-difference scores). To obtain a measure to compare both strategies, we calculated sample sizes needed in both models to obtain sufficient power to detect the association with increased schizophrenia risk.
**Figure 1.** Flowcharts of participants with 22q11DS for inclusion in the current study.

---

**TOTAL SAMPLE**

N = 1789
Individuals with a confirmed 22q11.2 deletion

**SAMPLE USED FOR NORMATIVE CHART a**
(Inclusion criteria: age between 6-38 b, W-test)

N = 1365
Total number of IQ measures = 2512

N = 657
with 1 IQ assessment

*Baseline*

Median age = 15.0
N SZ = 142 (21.6%)
N Males = 313 (47.6%)

N = 708
with 2 or more IQ assessments

*Longitudinal c*

Median age (at first assessment) = 10.4
N SZ = 101 (14.3%)
N Males = 354 (50%)

---

*a Data (cross-sectional and longitudinal) from these 1365 individuals were used for the construction of the normative chart (Methods section 2.2.i)*

*b For the current study we limited the age range to 6-38 years, as the main purpose of the study is to create one easily applicable normative chart. Above the age of 38 years, the number of participants in each age-year was too small (n < 10) to obtain a reliable normative value for that particular year. Below the age of 6 years, the number of participants in each age-year was also small and scores showed disproportionally greater variability (consistent with greater testing effects observed in IQ-tests in younger children).*

*c Data (longitudinal) from these 708 individuals were used for calculating the delta-z-scores (Methods section 2.2.ii)*
3. RESULTS

3.1 Participants and instruments

Figure 1 provides a schematic depiction of the participants included in this study. Table 1 provides descriptives for all participants, as well as separately for those with only one IQ-assessment (baseline) and those with two or more IQ-assessments available (longitudinal). Importantly, there were no differences in mean FSIQ, VIQ and PIQ scores (on the first available assessment) between the baseline- and longitudinal-samples (Table 1).

3.2.1 Normative chart statistics

The 3rd order polynomial regression provided the best fit for the FSIQ, VIQ and PIQ data, as indicated by the AIC and BIC, and the normative charts were constructed based on this. The parameters for the model for FSIQ were \( R^2 = 0.03, F(3,2508)=24.01, p<0.001 \); for VIQ \( R^2 = 0.03, F(3,2439)=19.19, p<0.001 \); and for PIQ \( R^2 = 0.03, F(3,2336)=26.35, p<0.001 \). Supplemental table 1 provides the coefficients of the regressions. The residuals of the model were normally distributed and constant over the age range, indicating accurate prediction of the trajectory by the normative chart. Further, the distribution of the Z-scores confirmed that the normative chart provided a good fit for the data. The mean Z-scores were close to 0 for FSIQ (-0.03), VIQ (-0.02) and PIQ (-0.03), indicating that on average, individuals with 22q11DS had an IQ-score as predicted by the model considering their age. In addition, there was no difference of the mean Z-scores between the baseline- and the longitudinal-samples. The standard deviations (SD) of the z-scores were close to 1 for FSIQ (1.02), VIQ (1.01) and PIQ (1.04)
(an SD of 1 is the equivalent of 15 IQ-points). **Figure 2** displays the normative growth chart for FSIQ, VIQ, and PIQ (including data points in **Supplemental Figure 1**).

In addition, as an illustration to aid in understanding the IQ decline observed on average in individuals with 22q11DS, **Supplemental Figure 2** represents the approximate corresponding raw score trajectory for 22q11DS, compared to raw IQ score change in the general population.

**Figure 2.** 22q11DS-specific normative chart for FSIQ (A), VIQ (B), and PIQ (C) over time.

These figures represent the normative charts for IQ development in individuals with the 22q11.2 deletion (A: FSIQ, B: VIQ, C: PIQ). The lines represent the observed average IQ trajectories (“Mean”), and the observed trajectories that deviate +/- 1 or 2 SDs from the mean. The norm charts are derived from 2512 IQ assessments in 1365 individuals with the 22q11.2 deletion between the ages of 6 and 38 years.
3.2.ii Delta Z-scores statistics

For the 708 individuals with longitudinal IQ-data were, we calculated delta-Z-scores; i.e., the difference between the Z-scores corresponding to the first and last available IQ-measurements. A model with a good fit would be expected to result in mean delta-Z-scores of around 0, as this would indicate that, on average, individuals stay on their trajectory. **Supplemental Figure 3** displays the distribution of the delta-Z-scores for FSIQ, VIQ and PIQ. The means were close to 0 (0.064, 0.069, and 0.089 respectively) and the standard deviations were 0.637, 0.679, and 0.720 respectively. Of the 708 individuals, 58% (FSIQ and VIQ) and 55% (PIQ) were between -0.5 and 0.5 SD. This indicates that on average, individuals stay on their trajectories as predicted by the normative IQ charts. **Figure 3**, presenting IQ data of two hypothetical individuals, serves to illustrate the enhanced impact of using delta-Z-IQ-scores (referenced to 22q11DS-specific norms) compared to general-population delta-IQ-values.

**Supplemental Materials 1** provides the calculator which allows for obtaining the expected IQ-score given a certain age, and hence the corresponding Z-score for an individual given their age and observed IQ-score. When multiple IQ-assessments for one individual are available, the delta-Z-scores can be calculated. This can be done for FSIQ, VIQ, and PIQ.
Figure 3. Two hypothetical cases that illustrate the advantage of 22q11DS-specific normative IQ-data over only (general population-based) IQ-data.

<table>
<thead>
<tr>
<th></th>
<th>Patient 1</th>
<th>Patient 2</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>First IQ assessment</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Age:</td>
<td>6 y</td>
<td>12 y</td>
</tr>
<tr>
<td>IQ:</td>
<td>75</td>
<td>75</td>
</tr>
<tr>
<td>Z-score:</td>
<td>-0.4</td>
<td>0.2</td>
</tr>
<tr>
<td><strong>Second IQ assessment</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Age:</td>
<td>12 y</td>
<td>18 y</td>
</tr>
<tr>
<td>IQ:</td>
<td>67</td>
<td>67</td>
</tr>
<tr>
<td>Z-score:</td>
<td>-0.4</td>
<td>-0.3</td>
</tr>
<tr>
<td><strong>Difference (between first and second assessment)</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Δ age:</td>
<td>6 y</td>
<td>6 y</td>
</tr>
<tr>
<td>Δ IQ:</td>
<td>-8</td>
<td>-8</td>
</tr>
<tr>
<td>Δ Z-score:</td>
<td>0</td>
<td>-0.5</td>
</tr>
</tbody>
</table>

Both patients show a difference of 8 IQ-points, over the timespan of 6 years (Δ IQ = -8). The patients were assessed at different ages. Using the 22q11DS-specific normative data results in different Z-scores corresponding to the patients' IQ-scores, given their age. Hence, the resulting Δ Z-scores are different between the patients, even though their Δ IQs are identical. Using 22q11-specific normative IQ data allows for detecting an important difference with clinical implications between the two patients, who show no differences when looking only at absolute IQ-scores: The decline of 8 IQ-points over 6 years in patient 1 is in keeping with the 22q11DS-specific IQ-trajectory (i.e., patient 1 remains on their IQ-curve; Δ Z-score = 0). The same decline in patient 2 however, exceeds the decline to be expected (i.e., patient 2 deviates from their IQ-curve; Δ Z-score = -0.5). Increased clinical monitoring may therefore be warranted for patient 2.
3.2.iii Post-hoc analyses

Post-hoc, we compared the Z-IQ-scores model to the population-based IQ model in terms of required sample size to detect the previously reported association between IQ-decline and schizophrenia risk [25]. As expected, both models reveal a significant association between VIQ-decline and schizophrenia. Importantly, the OR derived from the Z-based regression model was larger compared to the IQ-based model (Z-based OR = 2.84, 95% CI = 1.595 – 5.025, p = 3.405e-04; versus IQ-based OR = 2.09, 95% CI = 1.231 – 3.533, p = 5.862e-03). Based on these OR’s we calculated sample sizes needed to obtain sufficient power to detect the association between IQ-decline and schizophrenia. To obtain 80% power, the Z-based model required a sample size of 64, while a sample size of 91 was needed using the untransformed IQ values (Figure 4). In other words, using the Z-scores-based approach resulted in a 30% decrease of needed sample size, as compared to the untransformed IQ-based approach, to detect the association between IQ-decline and schizophrenia illness expression with 80% power.

Figure 4. Sample sizes of individuals with 22q11DS required to detect a significant effect (between VIQ-decline and schizophrenia) with 80% power: Z-score based approach versus IQ-based approach.
4. DISCUSSION

In this study we constructed a variant-specific normative chart for cognitive development from the largest sample of individuals with the 22q11.2 deletion available to date. Our findings suggest that in this population, a variant-specific normative IQ-chart can be reliably constructed and our discussion offers the rationale for how other pathogenic variants may benefit from a similar strategy. We propose that this approach allows for more accurate and informative interpretation of individual IQ-scores and trajectories, compared to using (untransformed) population-based IQ norms.

The findings further demonstrate that using variant-specific normative IQ data can significantly reduce the sample size needed to detect a certain effect (i.e., VIQ-decline and schizophrenia risk in 22q11DS), compared to population-based normative IQ data. From a research perspective, this is an important discovery. It is challenging to assemble adequately large datasets to provide sufficient power for phenotype-phenotype, or genotype-phenotype analyses, in particular with respect to longitudinal (deep-phenotyping) data. In populations with high-impact variants associated with neurodevelopmental outcome this challenge is even further magnified, given the low population-wide prevalence rates of such conditions.

22q11DS, IQ, and schizophrenia

Using data from 1365 individuals with the 22q11.2 deletion, our findings corroborate several important observations regarding IQ in this population. First, the data confirm the previously reported lower baseline IQ in individuals with 22q11DS [41,42], and show that the deletion thus appears to shift the IQ-distribution in carriers to the left (~-2 SD) as compared to non-carriers, but does not alter the characteristics of the
distribution. This is in line with a recent study which reported that while FSIQ, VIQ, and PIQ were ~30 IQ-points lower in 22q11DS patients compared to their unaffected parents, the distribution was normal and significantly associated to the parental distribution [submitted manuscript Fiksinski et al.].

Second, our data reiterate that in individuals with 22q11DS there is, on average, a decline in IQ over the lifespan [25] [23]. This observation underscores the impetus for regular and comprehensive cognitive assessments in individuals with 22q11DS [43-45]. We posit that in childhood and adolescence, the observed typical decline in 22q11DS mostly reflects a slower pace in cognitive development in individuals with 22q11DS, compared to their typically developing peers [46] (see also Supplemental Figure 2). In adulthood, however, this decline suggests that individuals with 22q11DS are losing cognitive capacities at a faster pace compared to the general population [31]. Importantly, using the 22q11DS-specific normative IQ-data allows for plotting an individual’s IQ-score or IQ-trajectory against expectations for this specific population.

Third, as previously reported [25], individuals with 22q11DS who show a VIQ decline that is steeper than what is expected based on the variant-specific trajectory are at a further increased risk for subsequently developing a psychotic disorder. This is in contrast to individuals with 22q11DS who do not deviate from their expected trajectory, but may still show a VIQ decline when compared to general population norms. These findings corroborate longitudinal studies in the general population, which report that individuals who later developed a psychotic disorder showed increasing cognitive impairments over time, especially during adolescence. In individuals from the general population who are at high risk for psychotic disorders, similar but milder delays in cognitive trajectories have been reported [47,48] [13,49,50].
Implications for this and other pathogenic variants

The often atypical and complex cognitive profile in carriers of pathogenic variants, such as the 22q11.2 deletion, adds complexity to the challenge of finding equilibrium between an individual’s profile of strengths and weaknesses on the one hand, and environmental demands on the other [43]. Realistic daily-life expectations given an individual’s capabilities are key in optimizing the fit between their individual profile and demands, and this is particularly important in populations with increased neurodevelopmental and psychiatric vulnerability [43,51].

Variant-specific normative IQ-data allow for “plotting” an individual’s IQ-score against norms given their specific variant, and, by extension, a likely projection into future performance. In other words, they allow for setting more realistic expectations and more informative monitoring of individual carriers of a pathogenic variant. For example, our data suggest that for a child with 22q11DS, a decline of 7 IQ-points between the ages of 7 and 13 is not unlikely. This is relevant in and of itself to the child’s day-to-day functioning with respect to environmental demands such as in school. The main relevance of knowing that this observed IQ-decline is in keeping with expectations given that this child has the 22q11.2 deletion is twofold. First, it aids in setting more realistic expectations in terms of future functioning and, potentially, taking proactive measures accordingly. Second, it may help to avoid unnecessary concern (e.g., in parents, teachers) as in fact, this child’s cognitive development is in line with the norms given their genetic variant and overall skills may not necessarily be deteriorating; but, rather, just demonstrating a slower rate of growth that is in line with the phenotypic performance for the condition.
Further, variant-specific normative IQ data enable the identification of those individuals who deviate more than what can be expected given the genetic variant; i.e., who deviate from their (adjusted) curve. This may be helpful in interpreting the observed IQ-decline and distinguishing between individuals who cannot keep up with increasing environmental (social, academic) expectations, and those who display an actual loss of abilities. While in both scenarios a decline in absolute IQ-scores can be observed, the underlying mechanisms and clinical implications may be very different [17,23,25,52]. Future studies could include raw IQ data (i.e., not standardized and norm-referenced) to further elucidate these different underlying mechanisms of IQ-decline, as well as to allow for further improved specificity and greater variance at the extremes end of the IQ-distribution [53].

Variant-specific normative IQ data may also allow for improved risk stratification for comorbid conditions. This strategy applied to 22q11DS facilitates the identification of those individuals with 22q11DS with a VIQ-decline in excess of what it typical for this population and that may be a significant risk factor for developing schizophrenia [25]. The clinical implication is that increased (early) monitoring for signs of psychotic development may be warranted in this subgroup. Vice versa, while still at increased risk of psychosis compared to the general population, the individuals who do not deviate from their expected trajectory (but may still show an IQ decline when compared to general population norms) could receive care as usual for 22q11DS patients [44,45]. Also, the stress experienced by patients and caregivers due to this genetically determined a priori risk for schizophrenia [54] may be somewhat mitigated in this group.
As is the case for 22q11DS, the variability in (degree of) expressed phenotypes with any rare pathogenic variant can still only be described in terms of group prevalence rates. Our current inability to provide individualized outcome prediction causes uncertainty for caregivers with respect to individual needs and daily life expectations [55,56], and undermines the potential for prevention or early intervention strategies. Although variant-specific normative data for IQ provide an important step towards improved outcome prediction at a group level, the identification of factors influencing individualized outcome prediction is needed. Recent studies are making progress in this regard in carriers of various high-impact genetic variants including 22q11DS, for example by investigating the impact of parental functioning on patient functioning on several phenotypes [57,58].

Similarly, more research is needed to further improve individualized risk stratification with respect to comorbid conditions and, subsequently, to elucidate how to potentially implement this in clinical practice. A recent IBBCC study shows promising progress in this area by demonstrating that the use of polygenic scores, in the context of a population with an a priori increased risk (22q11DS), can significantly improve the positive predictive value with respect to a particular phenotype; in this case schizophrenia [59].

**Strengths and limitations**

The main strength of this study is that we used IQ-data from the largest database of individuals with 22q11DS currently available. The multi-site collected data underwent extensive quality control, as described elsewhere [26]. We provide an easy-to-use normative IQ chart for the three main IQ constructs, which is readily accessible both to the clinical and research communities.
Limitations are that the available data did not allow for using independent samples in the two main parts of the analyses: creating the normative IQ chart (using all available data), and calculating the z-scores (using only longitudinal data), which would have been methodologically preferable. The results, however, provided confidence that our data were unbiased and indeed normative. Importantly, the data revealed no differences in IQ parameters between the subsets with longitudinal data available and the subset with only cross-sectional IQ-data (see also Table 1).

Our normative IQ-chart is limited to individuals with the 22q11.2 deletion between the ages of 6 and 38 (see also Footnote Figure 1), and the sample was not stratified for other key variables typically used in the development of normative tables (e.g., socioeconomic status, region of country). Future studies could include both younger and older individuals to expand coverage of the normchart to the entire lifespan of individuals with 22q11DS.

Finally, it is important to note that there are four key components of overall IQ that formally or informally permeate all versions of the Wechsler scales. Working Memory and Processing Speed are assessed independently from VIQ and PIQ and reflect key neuropsychological processes. Specific abnormalities in these domains may be associated with specific psychiatric or neurodevelopmental outcomes [31]. At the time of the current study, available data were limited to VIQ and PIQ, in addition to FSIQ. However, future studies that aim to elucidate Working Memory and Processing Speed data and trajectories in individuals with 22q11DS are important to further our understanding of the complete cognitive profile in individuals with this high-impact variant.
Conclusion

Here, we have discussed the rationale and methodology for using a normative chart for IQ and IQ development specific to a population with a specific pathogenic variant. Using the 22q11.2 deletion as a model, we demonstrate that a variant-specific IQ normative chart can be reliably constructed and offers important advantages over using only standard (general population) IQ norms. It allows for more informed interpretation and monitoring of cognitive performance in carriers of the pathogenic variant. It also contributes to the identification of individuals who deviate from their expected trajectory and may be at increased risk for clinically relevant comorbid conditions; e.g., in individuals with 22q11DS and a VIQ-decline steeper than what is expected in this population, the risk for schizophrenia is further elevated. We also demonstrated that using variant-specific normative IQ-data significantly reduces required sample size to detect relevant effects in a research context. The development of this normative chart, based on the largest sample of individuals with 22q11.2DS in the world, should provide additional opportunities to study the cognitive phenotypic presentation of this population specifically, but also provides a proof of principle regarding the identification of cognitive developmental trajectories in groups of individuals affected by other pathogenic variants. We expect that such knowledge will be valuable for clinical researchers and, ultimately, facilitate advances in clinical practice for these individuals and their families.
FUNDING AND DISCLOSURES:

Financial support: This study was supported in part by the NIMH International Consortium on Brain and Behavior in 22q11.2 Deletion Syndrome (U01MH101719 to RG); the Ter Meulen Grant of the Royal Netherlands Academy of Arts and Sciences and the UMCU Strategic Network Development Grant (AF); and the Brain and Behavior Research Foundation (formerly NARSAD; Young Investigator Award to JV).

Conflict of interest: All authors, including the members of the 22q11DS International Consortium on Brain and Behavior, declare that there are no conflicts of interest in relation to the subject of this study.

ACKNOWLEDGEMENTS:
The authors wish to thank all participants and their families for their participation in this study.

AUTHOR CONTRIBUTIONS:
Authors A.M.F., J.A.S.V., and E.J.B. had full access to the data and take responsibility for the accuracy of the data and analyses. Authors J.A.S.V. and E.J.B. contributed equally to this manuscript. All authors, including the members of the 22q11DS International Consortium on Brain and Behavior, contributed important intellectual content and reviewed the manuscript. A complete list of contributing authors, including the members of the 22q11DS International Consortium on Brain and Behavior, is listed in Supplemental Materials 2.
**TABLES AND FIGURE CAPTIONS/LEGENDS**

**Table 1.** Sample descriptives for total sample (n = 1789) of individuals with a 22q11.2 deletion, baseline subset (n = 657) and longitudinal subset (n = 708).

<table>
<thead>
<tr>
<th></th>
<th>Total sample: N = 1789</th>
<th>Subset: baseline: N = 657</th>
<th>Subset: longitudinal: N = 708</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Age in years at first assessment</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mean (SD)</td>
<td>17.1 (8.0)</td>
<td>15.0 (6.0 - 37.8)</td>
<td>11.6 (4.8)</td>
</tr>
<tr>
<td>Median (Range)</td>
<td>15.0 (6.0 - 37.8)</td>
<td>10.4 (6.0 - 35.2)</td>
<td></td>
</tr>
<tr>
<td><strong>Age in years at last assessment</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mean (SD)</td>
<td>17.9 (5.8)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Median (Range)</td>
<td>17.1 (7.4 - 38)</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Sex (% males)</strong></td>
<td>868 (48.5%)</td>
<td>313 (47.6%)</td>
<td>354 (50%)</td>
</tr>
<tr>
<td><strong>Psychotic illness expression</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Psychotic illness</td>
<td>332 (18.6%)</td>
<td>142 (21.6%)</td>
<td>101 (14.3%)</td>
</tr>
<tr>
<td>Control (age &gt;25 y)</td>
<td>295 (16.5%)</td>
<td>99 (15.1%)</td>
<td>63 (8.9%)</td>
</tr>
<tr>
<td>Putative control</td>
<td>850 (47.5%)</td>
<td>323 (49.2%)</td>
<td>385 (54.4%)</td>
</tr>
<tr>
<td>Control combined</td>
<td>1145 (64%)</td>
<td>422 (64.2%)</td>
<td>448 (63.3%)</td>
</tr>
<tr>
<td>Putative subthreshold</td>
<td>268 (15%)</td>
<td>74 (11.3%)</td>
<td>146 (20.6%)</td>
</tr>
<tr>
<td>Affective psychosis</td>
<td>33 (1.8%)</td>
<td>14 (2.1%)</td>
<td>13 (1.8%)</td>
</tr>
<tr>
<td>Unknown</td>
<td>11 (0.6%)</td>
<td>5 (0.8%)</td>
<td>0</td>
</tr>
<tr>
<td><strong>Age in years at last psychiatric assessment</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mean (SD)</td>
<td>21.3 (11.4)</td>
<td>19.9 (10.0)</td>
<td>19.0 (6.8)</td>
</tr>
<tr>
<td>Median (Range)</td>
<td>18.0 (2 – 71)</td>
<td>17.0 (5 – 56)</td>
<td>18.0 (7 – 61)</td>
</tr>
<tr>
<td><strong>IQ-test used (first assessment)</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>WPPSI</td>
<td>-</td>
<td>49 (7.5%)</td>
<td>32 (4.5%)</td>
</tr>
<tr>
<td>WPPSI-R</td>
<td>-</td>
<td>5 (0.8%)</td>
<td>18 (2.5%)</td>
</tr>
<tr>
<td>WISC-III</td>
<td>-</td>
<td>153 (23.3%)</td>
<td>323 (45.6%)</td>
</tr>
<tr>
<td>WISC-IV</td>
<td>-</td>
<td>96 (11.6%)</td>
<td>100 (14.1%)</td>
</tr>
<tr>
<td>WISC-R</td>
<td>-</td>
<td>24 (3.7%)</td>
<td>56 (7.9%)</td>
</tr>
<tr>
<td>WAIS-III</td>
<td>-</td>
<td>139 (21.2%)</td>
<td>49 (6.9%)</td>
</tr>
<tr>
<td>WAIS-IV</td>
<td>-</td>
<td>50 (7.6%)</td>
<td>9 (1.3%)</td>
</tr>
<tr>
<td>WASI</td>
<td>-</td>
<td>61 (9.3%)</td>
<td>7 (1.0%)</td>
</tr>
</tbody>
</table>

|                          |                          |                           |                               |
| **p**                    |                          |                           |                               |
| Mean baseline FSIQ (SD)   | 72.0 (14.3)              | 73.3 (13.1)               |                               |
| Mean baseline VIQ (SD)    | 76.3 (14.5)              | 76.9 (14.6)               |                               |
| Mean baseline PIQ (SD)    | 73.2 (14.9)              | 74.0 (13.4)               |                               |

*p-value of difference statistic (t-test) between baseline and longitudinal subsets.
Figure 1. Flowcharts of participants with 22q11DS for inclusion in the current study.

a Data (cross-sectional and longitudinal) from these 1365 individuals were used for the construction of the normative chart (Methods section 2.2.i)

b For the current study we limited the age range to 6-38 years, as the main purpose of the study is to create one easily applicable normative chart. Above the age of 38 years, the number of participants in each age-year was too small (n < 10) to obtain a reliable normative value for that particular year. Below the age of 6 years, the number of participants in each age-year was also small and scores showed disproportionally greater variability (consistent with greater testing effects observed in IQ-tests in younger children).

c Data (longitudinal) from these 708 individuals were used for calculating the delta-z-scores (Methods section 2.2.ii)

Figure 2. 22q11DS-specific normative chart for FSIQ (A), VIQ (B), and PIQ (C) over time.

These figures represent the normative charts for IQ development in individuals with the 22q11.2 deletion (A: FSIQ, B: VIQ, C: PIQ). The lines represent the observed average IQ trajectories (“Mean”), and the observed trajectories that deviate +/- 1 or 2 SDs from the mean. The normcharts are derived from 2512 IQ assessments in 1365 individuals with the 22q11.2 deletion between the ages of 6 and 38 years.

Figure 3. Two hypothetical cases that illustrate the advantage of 22q11DS-specific normative IQ-data over only (general population-based) IQ-data.

Figure 4. Sample sizes of individuals with 22q11DS required to detect a significant effect (between VIQ-decline and schizophrenia) with 80% power: Z-score based approach versus IQ-based approach.
REFERENCES


29 Wechsler D. Wechsler Adult Intelligence Scale-III. The Psychological Corporation: San Antonio, TX; 1997.
Kay SR, Fiszbein A, Opler LA. The positive and negative syndrome scale (PANSS) for schizophrenia. Schizophr Bull. 1987;13(2):261-76.


