The neurobehavioral phenotype in mucopolysaccharidosis Type IIIB: An exploratory study

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A B S T R A C T

Objective: Our goal was to describe the neurobehavioral phenotype in mucopolysaccharidosis Type IIIB (MPS IIIB). Parents report that behavioral abnormalities are a major problem in MPS III posing serious challenges to parenting and quality-of-life for both patient and parent. Our previous research on MPS IIIA identified autistic symptoms, and a Klüver-Bucy-type syndrome as indicated by reduced startle and loss of fear associated with amygdala atrophy. We hypothesized that MPS IIIB would manifest similar attributes when assessed with the same neurobehavioral protocol.

Methods: Ten patients with MPS IIIB were compared with 9 MPS IIIA patients, all older than 6. 8 younger children with Hurler syndrome (1H) were chosen as a comparison group for the Risk Room procedure; MPS IH does not directly affect social/emotional function and these younger children were closer to the developmental level of the MPS IIIB group. To examine disease severity, cognitive ability was assessed. Four evaluations were used: the Risk Room procedure (to measure social-emotional characteristics, especially fear and startle responses), the Autism Diagnostic Observation Schedule (ADOS), the Sanfilippo Behavior Rating Scale (SBRS), and amygdala brain volumes calculated from manually-traced MRI images.

Results: The two groups are equivalent in severity and show severe cognitive impairment. On the ADOS, the MPS IIIB patients exhibited the same autistic features as IIIA. The IIIB means differed from MPS IH means on most measures. However, the IIIB group did not approach the Risk Room stranger, like the MPS IH group who kept their distance, but unlike the IIIA group who showed no fear of the stranger. On the SBRS, the MPS IIIB patients were described as more inattentive and more fearful, especially of new people than the MPS IIIA. Onsets of some disease characteristics appeared more closely spaced and slightly earlier in MPS IIIB than IIIA.

Conclusions: On most behavioral measures, MPS IIIB patients did not differ substantially from MPS IIIA patients over age six, demonstrating autistic features and a Klüver Bucy-like syndrome including lack of fear and poor attention. Delay in onset of behavioral symptoms was associated with later diagnosis in two patients. Lack of fear, poor attention, and autistic-like symptomatology are as characteristic of MPS IIIB as they are of MPS IIIA. A possible difference is that the some behavioral abnormalities develop more quickly in MPS IIIB. If this is so, these patients may become at risk for harm and present a challenge for parenting even earlier than do those with MPS IIIA. In future clinical trials of new treatments, especially with respect to quality of life and patient management, improvement of these behaviors will be an essential goal. Because very young patients were not studied, prospective natural history documentation of the early development of abnormal behaviors in MPS IIIB is needed.

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1. Introduction

Mucopolysaccharidosis type IIIB (MPS IIIB) is an autosomal recessive lysosomal disorder associated with severe behavioral abnormalities and progressive dementia. Of the four subtypes of MPS III, it is reported to be the second most common (approximately 1 in 200,000 births) [1,2] with the exception of south-eastern European countries, where it may be the most common of MPS III types [3–5].

It is caused by reduced N-acetyl-alpha-D-glucosaminidase (NAGLU; EC 3.2.1.50) catalytic activity, a necessary metabolic step in degradation of the glycosaminoglycan heparan sulfate, leading to accumulation of heparan sulfate and secondary gangliosides in the central nervous system [6,7].
Behavioral abnormalities in MPS IIIB have been noted but not described in detail [8,9]. We have previously reported that, in the classic, severe form of MPS IIIA, many characteristics of autism begin to appear at about 3–4 years of age [10]. We also found that the behaviors in MPS IIIA, including increased orality, reduced startsle, and reduced fearfulness, present as a variant of Klüver-Bucy syndrome [11,12]. We also found a significantly greater rate of amygdala atrophy over time associated with the reduced fearfulness [11].

In the present study, we hypothesized that MPS IIIB would display behavioral abnormalities and rate of appearance of behavioral symptoms similar to those of MPS IIIA. If similar patterns do emerge, it could provide guidance for understanding the natural history of the disease and markers for treatment response. If not, expectations for disease and treatment response will need to be altered for MPS IIIB.

2. Methods

2.1. Participants

2.1.1. MPS IIIB patients

During the recruitment phase, due to low incidence of MPS IIIB, accrual was considerably lower than anticipated, so we were unable to recruit younger and less impaired children. Thus, our findings are restricted to older patients with more advanced disease.

Parents of 11 children with MPS IIIB, ages 0.5 to 32 years, were contacted for this cross-sectional neurobehavioral study from a longitudinal natural history (NH) study sponsored by Shire at the University of Minnesota (NCT01508768). The goal of the longitudinal study was to detail the natural course of MPS IIIB and to identify potential endpoints for future treatment trials.

Patients in the NH study met the following criteria: (a) MPS IIIB diagnosis confirmed by enzyme or genetic assay, (b) developmental age ≥12 months on the Vineland Adaptive Behavior Scales, Second Edition [13]. Exclusion criteria were: (a) known hyper-sensitivity to anesthesia, (b) history of ameliorative treatment with any investigational drug or device precluding MRI, and (g) blindness and/or deafness.

Inclusion criteria for this neurobehavioral study were being more than two years old and independently ambulatory, which excluded one child from the NH study who was less than one year old for a total sample size of 10. Although the NH study had European sites as well, the neurobehavioral study was only carried out at the baseline visit to the University of Minnesota.

2.1.2. MPS IIIA comparison group

From a previous natural history study of MPS IIIA (NCT01047306), we matched our sample to 9 patients in a similar age range, all over the age of six. This subsample was part of a larger sample of MPS IIIA whose behavioral data have been reported previously [10–12].

2.1.3. MPS IH comparison group

In addition, the 8 children with MPS I (Hurler syndrome, or MPS IH) who participated in the MPS IIIA neurobehavioral study were selected as a comparison group for this Risk Room procedure [11]. This younger group was selected as they have a similar level of cognitive development.

2.2. Procedures

2.2.1. Neurocognitive assessment

To determine the phenotypic severity and the equivalency of our two samples, cognitive ability scores were utilized. As part of the NH studies mentioned above, two standard instruments, the Bayley Scales of Infant Development, Edition III (BSID-III) [14] or the Kaufman Assessment Battery for Children, Edition II (KABC-II) [15] were selected that cover the anticipated age and ability range. Age equivalent scores (AEqs) were generated from published normative data [14,15] and a Development Quotient (DQ) was derived by dividing the AEq by chronological age and then multiplying by 100. This approach avoids the “floor” effects in normative tables that makes scores insensitive for severely cognitively impaired children [16–18]. The specific test administered was chosen according to an algorithm previously described [19].

2.2.2. Risk Room

The rationale for this laboratory procedure is outlined in our description for the MPS IIIA patients [11]. The Risk Room is a part of the Laboratory-Temperament Assessment Battery [20], which evaluates social/emotional behaviors, and specifically fearfulness, in lower functioning children [20]. To test whether the MPS IIIB patients presented with a similar variant of Klüver-Bucy syndrome and to investigate attachment/compliance issues, we assessed participants’ 1) locomotor exploration of the Risk Room, 2) response to attractive and mildly frightening objects in the room (scary Halloween masks and a seated male stranger wearing a ball cap and sunglasses), 3) exposure to a 92-dB startle-noise triggered by the patient’s first contact with an attractive toy (the “trigger toy”). This test is based on findings that amygdala dysfunction reduces startle in other animals [21,4] reunion with their mother after her brief absence, and 5) compliance with her directive to pick up and put away a small toy [22].

2.2.3. Autism Diagnostic Observation Schedule

The ADOS is a semistructured observation tool designed to observe and judge the quality of a child’s social communication and play and to assess for the presence of any excessively intense interests or repetitive behaviors [23]. The revised algorithms of the ADOS were used [24]. The ADOS yields total scores for social affect and restricted and repetitive behavior, yielding an overall classification indicating behaviors and symptoms consistent with autism, consistent with milder indications of autism spectrum disorder (ASD), or not consistent with ASD (“nonspectrum”).

2.2.4. Sanfilippo Behavior Rating Scale (SBRBS)

Parents completed the 68 items in the SBRBS [12]. Details of that measure can also be found at z.umn.edu/sbrbs. Four cluster scores and two domain scores are generated as well as parental recall of age of onset of behavioral symptoms. The four cluster scores are Movement, Lack of Fear, Social-Emotional, and Executive Function; a higher score denotes more abnormality in each cluster. Each cluster score contains a number of domains and individual items that make up the domain. Two domains are unique and not included in clusters: “Mood, Anger, and Aggression”, and “Orality” (excessive mouthing of objects).

2.2.5. Amygdala and hippocampus volumes

While the patient was under anesthesia during the baseline visit, MRI was acquired on a 3-Tesla Siemens Trio scanner with sequences that included MPRAGE (magnetization-prepared rapid acquisition with gradient echo). Volumetric analyses included manual tracing of amygdala and hippocampus usingBrains2, which allows accurate measurement of three-dimensional representations of structures [25]. Our rater’s intra-rater reliability was 0.99 and inter-rater reliability was 0.87 [26].

2.2.6. Statistical analyses

Group data were summarized by means and standard deviations for continuous variables and frequency with percentages for categorical variables. Confidence intervals for estimated means within a group were based on a t-distribution; differences in means were evaluated using a t-test with unequal variance and Welch approximation to the degrees of freedom. Confidence intervals for estimated proportions within a group were determined by inverting the score test; differences in proportions between groups were evaluated with chi-squared tests.
Survivor analytic techniques (time-to-event analysis) have been used to address the timing of symptom appearance in disease \[27\]. All analyses were conducted using R v3.1.1 \[28\].

### 3. Results

All MPS IIIB patients tested were Caucasian (as were the comparison groups). 6/10 MPS IIIB and 5/9 MPS IIIA patients were male. The average (SD) age for the MPS IIIB patients was 16.3 (9.5) years and for the MPS IIIA patients was 10.0 (4.5) years. Four patients in the IIIB group were over age 20 (ages 22, 23, 28, and 32); the oldest in the IIIA group was 18.4. We analyzed the data with and without the two patients in the IIIB group who were outliers in age, both female siblings, ages 28 and 32, diagnosed within the past two years. The average age at diagnosis was 89 months in the MPS IIIA group and 119 for the III B group. However with the two older patients removed, the average age at diagnosis for the MPS IIIB group was 64 months.

In order to determine whether the two groups differed in phenotypic severity, we report their age equivalent scores on the BSID or the non-verbal scale of the KABC-II. Results are reported in Table 1 with the outliers removed.

The two outliers in age have a chronological age of 27.77 and 31.70 with cognitive age equivalent scores of 21 and 10 months respectively. These low scores do not rule out the possibility they could have attenuated disease as their diagnosis was so late and no early testing was available. The single child who was not included because the age was below the inclusion for the risk room and ADOS, at 1.69 years had an age equivalent score of 17 months yielding a developmental quotient of 84. This is similar to the range of results of the MPS IIIA patients of the same age \[29\].

#### 3.1. ADOS

The results on the ADOS were very similar to those of the MPS IIIA group (Table 1). Of the 10 MPS IIIB patients, scores for all but one were in the ADOS range consistent with those on the autism spectrum (12 is the cutoff); higher scores indicate more behaviors associated with autism spectrum disorder). The one patient who did not meet ADOS criteria was 6.7 years old and had an ADOS total score of 9; a 6 year old participant had the next highest score exceeding the cutoff with a score of 16. All others had scores between 16 and 23 compared with the MPS IIIA patients whose range of scores was 12–22. The social/affective domain was more affected than the restricted or repetitive behaviors in both groups.

#### 3.2. Risk Room

#### 3.2.1. Comparison with MPS IH

As expected, the MPS IIIB patients differed significantly from the MPS IH comparison group on four of nine measures (Table 2). With the older two patients removed, three of nine measures were statistically significant and one indicated a trend. Although substantial mean differences were found, lack of significance on several measures is due to variability and small N in both groups.

### Table 1

| Age and cognitive ability (means (M), standard deviations (SD) and range) for MPS IIIB and MPS IIIA. |
|---|---|---|---|---|
| MPS IIIB (without outliers) | MPS IIIA |
| M (SD) | Range | M (SD) | Range |
| Age (years) | 12.94 (7.01) | 6.28–23.95 | 9.95 (4.46) | 6.41–18.33 |
| Cognitive age equivalent (months) | 16.25 (8.40) | 8–28 | 16.89 (12.56) | 6–45 |
| Developmental quotient\(^a\) | 14.75 (13.00) | 3–37 | 14.79 (7.79) | 3–26 |

\(^a\) Calculated using age equivalent score divided by chronological age (Delaney et al.).

### Table 2

ADOS Scores, mean (SD), for MPS IIIB (with outliers) and IIIA and difference in means.

| ADOS SOCIAL | ADOS BEHAVIORAL | ADOS TOTAL |
|---|---|---|
| MPS IIIB (n = 9) | MPS IIIA (n = 9) | Difference (95% CI) | P-value |
| 14.3 (4.4) | 15.6 (4.3) | 1.2 (−5.6, 3.1) | 0.56 |
| 3.8 (1.1) | 3.2 (1.2) | 0.6 (−0.6, 1.7) | 0.32 |
| 18.1 (4.3) | 18.8 (3.5) | −0.7 (−4.6, 3.3) | 0.72 |

\(^a\) One patient did not have an ADOS evaluation.

#### 3.2.2. Comparison with MPS IIIA

Only a few mean differences are noted overall (p values are in Table 3).

#### 3.2.3. Exploration

The MPS IIIB patients were similar to MPS IIIA and differed notably from the IH patients in exploration of the room and proximity to their mothers. The IIIB patients spent an average of 62% of their time walking around the room (53.8% with the oldest two patients removed), similar to 59.7% for IIIA, compared to 10% for the MPS IH group. The MPS IIIB patients did not differ significantly from the IIIA patients in time spent next to their mother (30.0% and 28.0%, respectively) compared to 71.5% for MPS IH. Similarly 70.0% of the IIIB patients touched a scary mask at least once compared to 77.8% for IIIA and none for the MPS IH group. Removing the older patients did not change the significance values for time next to mother or touching the mask. However, fewer MPS IIIB patients (30%) approached the stranger than did the MPS IIIA group (37.5% with older patients removed) compared to 66.7 although the differences were not statistically significant. The MPS IIIB was similar to the MPS IH group, only 33.3% of whom approached.

#### 3.2.4. Response to startling noise and trigger toy

The proportion of the MPS IIIB patients who startled to the loud noise (30%) was similar to that of the previously tested the IIIA group (33.3%), but significantly less than the 75.0% of the MPS IH patients who startled. Sixty percent of the IIIB group returned to the trigger-toy following the noise compared to 78% of the IIIA group. 12.5% of the MPS IH group returned to the trigger-toy. Removing the two older patients did not substantially change these results.

#### 3.2.5. Reunion score and compliance

During the 1–2 min following the mother’s return from her brief exit, the IH group showed the most response to her, the IIIA group showed less response while the IIIB patients acknowledged her the least. When asked to pick up and put away a toy, the IIIB group (40%) was more compliant than the IIIA group (22.2%), but not as compliant as the IH group (87.5%). With the two older patients removed, 50% percent of the IIIB group demonstrated compliance.

#### 3.3. SBRS

#### 3.3.1. Level of behavioral impairment

The MPS IIIB group was similar to the IIIA group in the frequency of items endorsed on three of the four clusters: Movement, Lack of Fear, and Executive Function (Table 4). On the Social-Emotional Cluster, the IIIB patients tended to score higher than IIIA patients, although this trend was nonsignificant. The two MPS III groups did not differ from each other on the two domains, Orality and Mood/Anger/Aggression. We examined the domains within each cluster, and found that within the Lack of Fear Cluster, the Safety Consciousness domain overall was equally impaired in the IIIB group (with or without the older patients) and the IIIA groups. Within the Social-Emotional Cluster, Emotional Functioning was slightly more impaired in the IIIB group with and without the older patients suggesting more emotional disruption. Within the Executive Function Cluster, the Attention domain was significantly

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more impaired in the IIIB group. After removing the older two patients, the means remained the same but a statistically significant difference shifted to a trend.

Parent ratings for individual items indicated that IIIB patients were significantly less likely to go into unsafe areas and scored lower for lack of fear than did IIIA patients, but IIIB patients were described as slightly more fearful of new people. Again a trend was present for attention to the means remaining the same but a statistically signi-

3.3.2 Ages of symptom onsets

When parents recalled the age of onset of behavioral symptoms, the IIIB patients were described as having a later onset on average than IIIA patients in almost every domain (Table 5). However, when the two older patients were removed, this difference disappeared. Data for 6 MPS IIIB patients (without the oldest two) and 7 MPS IIIA patients were available for a time-to-event analysis of age of onset. Fig 1 plots the successive ages of onset of each behavior characteristic within each group and compares the relative rate of appearance of each characteristic between the two groups. The time-to-event intervals for Executive function and Mood/anger/aggression were shorter within the IIIB group than within the IIIA group. Lack of Fear showed a trend in the same direction.

Comparison of MPS IIIA and IIIB within-group time-to-event plots for onsets of six behavior clusters and domains. The same age scale is used in all plots. Final sample proportions are <1.0 due to missing data. Between group differences in the right hand plots show a trend or are significant.

3.4 Amygdala and hippocampus volumes

No meaningful or statistical differences were found in baseline amygdala and hippocampus volumes between IIIB and IIIA patients (Table 6).

Table 3

Mean/percent with 95% confidence intervals of risk room scores across groups.

| Variable                      | MPS IIIB (N = 10) | MPS IIIA (N = 9) | P-value A vs B | P-value IH vs. B | MPS III B (N = 8) | P-value A vs B | P-value IH vs. B |
|-------------------------------|-------------------|-----------------|----------------|------------------|------------------|----------------|------------------|
| Move away latency (seconds)   | 12.0 (9.0, 15.0)  | 25.6 (15.4, 49.6) | 1.0 (0.0, 395.9) | 0.322            | 0.167            | 12.5 (8.6, 16.4) | 0.249 (0.168) |
| Percent time in locomotion    | 62.0 (38.7, 85.3) | 59.7 (33.2, 86.2) | 10.0 (0.0, 22.3) | 0.082            | <0.001           | 53.8 (27.7, 79.8) | 0.715 (0.005) |
| Percent time next to mother   | 27.4 (11.3, 43.5) | 28.0 (3.4, 52.6)  | 71.5 (41.71013) | 0.963            | 0.010            | 30.8 (10.7, 30.8) | 0.843 (0.018) |
| Touch mask (%)                | 70.0 (35.4, 91.9) | 77.8 (40.2, 96.1) | 0.0 (0.4, 84.3) | 1.000            | 0.027            | 62.5 (25.9, 88.9) | 0.875 (0.064) |
| Approach stranger (%)         | 30.0 (8.1, 64.6)  | 66.7 (30.9, 91.0) | 33.3 (6.0, 75.9) | 0.255            | 1.000            | 37.5 (10.2, 74.1) | 0.474 (1.000) |
| Startle (%)                   | 30.0 (8.1, 64.6)  | 33.3 (30.9, 69.1) | 75.0 (35.6, 95.5) | 1.000            | 0.155            | 37.5 (10.2, 74.1) | 1.000 (0.313) |
| Hold/return to trigger toy (%)| 60.0 (27.4, 86.3) | 77.8 (40.2, 96.1) | 12.6 (31.5, 53.3) | 0.735            | 0.117            | 62.5 (25.9, 88.9) | 0.875 (0.121) |
| Clean-up compliance (%)       | 40.0 (9.0, 2.5)   | 22.2 (13.9, 59.8) | 87.5 (46.7, 99.3) | 0.735            | 0.117            | 50.0 (21.5, 78.5) | 0.492 (0.281) |

Superscript in MPS IH column denotes number missing.

a Designates group with outliers (older patients) removed.
b 0 = ignore mother, 1 = look at mother, 2 = talk to mother without approaching, 3 = approach mother, 4 = make physical contact/hug.

Table 4

Results from SBRs (range of scores are from 1 to 6 with higher numbers being more abnormal) comparing results from MPS IIIB with MPS IIIA.

| Variable                  | MPS IIIB (N = 10) | MPS IIIA (N = 8) | P-value |
|---------------------------|-------------------|-----------------|---------|
| Cluster                   |                   |                 |         |
| Movement cluster          | 2.2 (0.7)         | 2.1 (0.9)       | 0.684   |
| Lack of fear cluster      | 2.3 (0.7)         | 2.7 (0.8)       | 0.222   |
| Social emotional cluster  | 2.4 (0.6)         | 1.9 (0.6)       | 0.151   |
| Executive function cluster| 3.6 (1.1)         | 3.4 (1.1)       | 0.656   |
| Unique domains (not in cluster) |         |                 |         |
| Mood, Anger, Aggression   | 1.5 (1.0)         | 2.1 (1.1)       | 0.223   |
| Orality                   | 2.5 (1.4)         | 3.1 (1.7)       | 0.432   |
| Domains of interest (within clusters) |         |                 |         |
| Safety Consciousness      | 3.1 (1.2)         | 3.9 (1.4)       | 0.223   |
| Emotional function        | 2.2 (0.9)         | 1.5 (0.9)       | 0.132   |
| Attention                 | 4.2 (0.7)         | 3.5 (0.6)       | 0.030   |
| Items of interest (in domains/cluster) |     |                 |         |
| 26. Goes into unsafe areas| 3.4 (1.7)         | 5.1 (1.2)       | 0.027   |
| 28. Less fearful           | 2.1 (1.5)         | 3.6 (1.6)       | 0.058   |
| 32. Fearful of new people  | 1.2 (1.5)         | 0.2 (0.7)       | 0.117   |
| 53. Does not pay attention | 3.9 (1.1)         | 2.9 (1.2)       | 0.095   |

Superscript denotes number missing.
a Designates group with 2 outliers (older patients) removed.
b One patient did not complete the SBRs.

Table 5

Age of onset of symptoms in years from SBRS ratings by parents.

| Variable                  | MPS IIIB (N = 10) | MPS IIIB (N = 8) | MPS IIIA (N = 8) | Mean (SD) age of onset | Mean (SD) age of onset | Mean (SD) age of onset |
|---------------------------|-------------------|-----------------|-----------------|------------------------|------------------------|------------------------|
| Cluster                   |                   |                 |                 |                        |                        |                        |
| Movement cluster          | 8.2 (6.9)         | 6.7 (2.0)       | 4.6 (3.8)       | 4.67 (2.03)            | 4.14 (3.00)            |                        |
| Social-emotional cluster  | 8.69 (8.12)       | 4.33 (2.63)     | 4.62 (3.19)     |                        |                        |                        |
| Lack of fear cluster      | 5.60 (4.46)       | 3.66 (1.73)     | 3.90 (2.52)     |                        |                        |                        |
| Executive function cluster| 5.26 (4.05)       | 3.89 (2.26)     | 4.56 (3.29)     |                        |                        |                        |
| Domains:                  |                   |                 |                 |                        |                        |                        |
| Mood, anger and aggression| 6.40 (5.17)       | 4.38 (1.36)     | 5.78 (4.08)     |                        |                        |                        |
| Orality                   | 10.64 (10.11)     | 5.08 (3.65)     | 6.43 (6.38)     |                        |                        |                        |
| Mean across all areas     | 7.47 (6.48)       | 4.33 (2.28)     | 4.91 (3.69)     |                        |                        |                        |

9.3% of missing values for age of onset in MPS IIIB and 21.2% missing values for MPS IIIA.
a Designates group with outliers (older patients) removed.
b Sample size is 8 due to one patient missing SBRS.
4. Discussion

MPS III is characterized by progressive neurodegeneration and dementia, with death typically occurring in the second decade of life [1]. In MPS IIIA, symptoms become apparent between 2 and 6 years of age, although diagnosis often lags behind the appearance of the earliest symptoms [30]. We have previously reported that MPS IIIA has a characteristic behavioral pattern consisting of autism-like symptoms and what appears to be a variant of Klüver-Bucy Syndrome including orality, reduced startle and fearfulness, and associated with atrophy in amygdala volume [10–12].

Our primary objectives were to determine how patients with MPS IIIB differ behaviorally from those with MPS IH and, more specifically, from those with IIIA, especially with regard to autistic and Klüver-Bucy symptoms. We were able to characterize the behavior of the MPS IIIB participants in our group; however, our recruited sample was older and showed more disease progression than we anticipated. We were able to select an older group of patients with MPS IIIA in a more comparable age range, which allowed us to compare behavioral symptoms after age 6 between these two disease groups. However, low incidence of MPS IIIB in the U.S. and low accrual in this study, limited our ability to characterize the behavioral course of the disease for patients under six.

In order to ensure that our patients did not have an attenuated phenotype and that they were equivalent in stage of disease, we compared them on measures of cognitive development. Both groups were profoundly cognitively impaired. The two outliers were similarly impaired, but as they were diagnosed later than any other patient in our samples, we did not include them in our comparison. Clearly compared to the patients described by Valstar [31], our sample did not fall into the attenuated group. We surmised that phenotypic cognitive data would be more

Table 6
Hippocampus and amygdala volumes in ml.

|                    | MPS IIIB (n = 9) | MPS IIIA (n = 9) |
|--------------------|-----------------|-----------------|
| Left hippocampus   | 1.83 (0.29)     | 1.84 (0.29)     |
| Right hippocampus  | 1.89 (0.23)     | 1.91 (0.26)     |
| Left amygdala      | 1.01 (0.11)     | 1.04 (0.23)     |
| Right amygdala     | 1.04 (0.09)     | 1.07 (0.24)     |

![Fig. 1. Age of onset data from SBRS as recalled by parents.](image-url)
of symptom onsets in the remaining MPS IIIB patients were compared to onsets in MPS IIIA (Fig. 1), some disease characteristics appeared to present more rapidly and occur slightly earlier across the IIIB group. Onsets were more rapid for Executive Function and Mood/anger/aggression, with a trend in that direction for Lack of Fear.

There is growing interest in the significance of age at onset of disease, e.g., disorders being more severe, persistent and less amenable to treatment when they appear earlier rather than later in childhood [37–39]. In some cases, earlier age of onset of the first disease symptom is associated with the earlier appearance of the full range of disease symptoms [38].

Note that for both groups, but especially MPS IIIA patients, some age of onset data were missing. Parents’ poor memory for behavior onset may be why they were reluctant to complete this portion of the questionnaire. While missing data may bias these results, which should certainly be viewed as exploratory, they do demonstrate the need for prospective natural history data in the youngest MPS IIIB patients to clarify ages of symptom onset.

Overall, our results suggest that in those over 6 years old, MPS IIIB patients, like MPS IIIA patients, do not fear scary objects but are slightly more fearful of strange people. However, differences in these behaviors presumably related to Klüver-Bucy syndrome are not reflected in their respective amygdala volumes, which are similar in both groups. We also have exploratory evidence of more attention difficulties in the IIIB patients.

Our two oldest patients demonstrate a different course that has a later onset and progresses more slowly, although it is ultimately as severe. Such patients are likely to be identified as having another disorder, such as autism or attention deficit hyperactivity disorder, before their cognitive decline is noticed. For this reason their diagnostic odyssey may be more protracted and locating these patients is more difficult. Misdiagnosis leading to delayed identification of MPS III disorders has been previously documented [40].

We conclude that behavioral patterns in MPS IIIB and IIIA patients over age 6 do differ in a few areas but both demonstrate symptoms associated with autism and Klüver-Bucy syndrome. However, because we were unable to recruit younger patients, unanswered questions remain about the onset and course of the behavioral presentation prior to age 6. In our MPS IIIB sample, and even in the oldest two patients symptom severity does not appear to be attenuated, unlike in the European sample previously described [31].

We caution that because this is an exploratory study, some mean differences in the results we describe lack statistical significance. In rare diseases, lack of power to detect significant effects is a frequent problem. A larger international sample using a multicenter approach is needed to verify these exploratory results. Clearly, longitudinal research is needed, particularly in parts of the world where MPS IIIB is more prevalent.

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