First episode psychosis with extrapyramidal signs prior to antipsychotic drug treatment

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Extrapyramidal movement disorders are common in chronic schizophrenia, and may be an intrinsic feature of the illness as well as related to antipsychotic drug treatment. Similar dysfunctions at illness onset may have implications for outcome, and for understanding the mechanisms of illness. The objectives were to examine the clinical correlates of pre-treatment movement disorders at first episode of psychosis, and determine associations with neuropsychological function and striatal structure. Never medicated subjects were recruited from consecutive admissions to Early psychosis Programs with defined catchment areas in Hong Kong, China, and Halifax, Canada. Standardized clinical, neuropsychological and brain imaging assessments were carried out at baseline and following acute and long term treatment with typical or atypical antipsychotic drugs. At the Hong Kong site, we studied 84 subjects with first episode psychosis (n = 10 with EPS). At the Halifax site, we studied 40 subjects with first episode psychosis (n = 17 with EPS), and 23 healthy comparison subjects. Subjects with movement disorders prior to treatment (EPS+) had higher total PANSS scores at baseline (mean elevation 19.9% Hong Kong, \( P = 0.016 \); 14.7% Halifax, \( P = 0.049 \)). In subjects treated with atypical antipsychotics (all Halifax), EPS+ status at baseline predicted more movement disorders at long term follow up (\( P = 0.0005 \)). In both cohorts, EPS+ subjects had poorer acute symptomatic treatment response assessed with the PANSS (Hong Kong \( P = 0.005 \); Halifax \( P = 0.017 \)). Neuropsychological impairment related to executive dysfunction appeared greater in a small sample of EPS+ subjects (Hong Kong, effect size 0.26–0.27, \( P < 0.05 \)). Caudate volumes were 4.5% larger in EPS+ compared with EPS-subjects (Halifax \( P = 0.042 \)), and correlations between striatal volumes and age were different in the EPS+ group. In conclusion, pre-treatment EPS is present in a substantial minority of subjects with first episode psychosis, appears to persist at long term follow up, and is associated with poorer response of symptoms to treatment. Selective impairment of executive function and striatal enlargement provides evidence of abnormalities of brain function and structure associated with this aspect of early psychosis.

schizophrenia, movement disorder, basal ganglia, antipsychotics, cognitive, psychosis

Signs of extrapyramidal dysfunction including akinesia, tremor, dystonia and dyskinesia were recognized in patients with schizophrenia prior to use of antipsychotic drug treatments [1–4]. Many of these same signs are described in family members of patients [5,6]. Most endophenotype studies investigate neurological soft signs, that appear to represent signs of developmental deviance [7,8]. Extrapyramidal dysfunction may link more closely to the neuropharmacological features of schizophrenia, but must be differentiated from the effects of treatments used for the illness. With the introduction of antipsychotic drugs in the 1950s, and particularly with the development of high affinity dopamine D2
antagonists, extrapyramidal signs became commonplace, complicating the investigation of movement disorders intrinsic to schizophrenia. However, implementation of early identification and intervention strategies for psychosis has reawakened interest in the movement disorders in schizophrenia, as has the search for endophenotypes of the illness.

Contemporary studies indicate acutely ill, never medicated patients may have signs of Parkinsonism (4%–28%), dystonia (2%–17%) and dyskinesia (1%–14%) [9–18]. Clinical correlates of extrapyramidal signs in first episode psychosis include increased negative symptoms with akinetic Parkinsonism in some [9,17,18], but not all studies [15]. Lower education was a predisposing feature for dyskinesia prior to treatment [11]. Recent studies of first episode patients report that poor premorbid functioning and possibly a family history of severe mental illness were associated with signs of extrapyramidal dysfunction prior to antipsychotic exposure [17,18].

Several lines of evidence, including a series of studies of older, chronic, never medicated patients suggest that movement disorders may be an integral part of schizophrenia [19,20]. Dyskinesia was the main manifestation in a chronic patient cohort, but Parkinsonism was also present [21,22]. Follow up studies over 18 months showed the movement disorders fluctuated over time, with 57% affected by dyskinesia and 35% affected by Parkinsonism on at least one assessment [22]. Dyskinesia was not related to memory dysfunction [23]. These patients showed enlargement of the left lentiform nucleus, and loss of the negative correlation between age and volumes of the caudate and lentiform nuclei [21].

Similarly detailed studies in early psychosis could be equally informative. The present investigation was designed to test the hypotheses that extrapyramidal signs at first presentation of psychosis are associated with clinically relevant features of illness, predict response of symptoms to treatment, and are associated with neuropsychological dysfunction and structural abnormalities of the striatum.

1 Subjects and methods

1.1 Subjects

Data was obtained from subjects treated by first episode psychosis programs in Hong Kong and in Halifax, and from healthy comparison subjects recruited in Halifax. All subjects provided written informed consent, and protocols were approved by the respective Ethics Committees. The Hong Kong sample was obtained from 153 consecutive first admissions for psychosis to Queen Mary and Pamela Youde Nethersole Eastern Hospitals, serving the Hong Kong Island catchment area (population 1300000). The time period was September, 1997 to March, 2000. The Halifax sample was obtained from approximately 250 consecutive referrals to the Nova Scotia Early Psychosis Program, which serves a Provincial population of 940000. The time period was December, 1995 to March, 2000.

Each program obtained standardized assessments of subjects at entry and after follow up. Assessments were made by program psychiatrists, trained in research methodology. Where possible, subjects were assessed prior to initiating antipsychotic treatment (Hong Kong n = 84, Halifax n = 40). The majority of subjects remained in treatment for at least 1 year. At both sites, inclusion criteria were: no previous antipsychotic drug treatment, a first episode psychotic disorder established according to DSM-IV criteria using all available clinical information after at least 6 months of assessment, age 13–65, in- or out-patients. Exclusion criteria were: known neurological disorder, or diagnosis of psychosis related to substance abuse. As part of the Hong Kong site protocol, subjects were administered neuropsychological tests. An additional inclusion criterion was fluency in Cantonese. Subjects with a history of special school attendance (usually indicating moderate to severe learning disability) were excluded. As part of the Halifax site protocol, subjects could participate in an MRI scan. Healthy comparison subjects were recruited from hospital staff and the local community. For the MRI study, exclusion criteria were a history of significant head injury or loss of consciousness greater than 5 min, a history of DSM-IV substance abuse or seizure disorder, and a family history of psychosis (for comparison subjects).

1.2 Assessment of extrapyramidal signs

Assessments were made prior to antipsychotic treatment. For the Hong Kong sample, subjects were assessed with the Simpson-Angus Scale (SAS) [24], and the Abnormal Involuntary Movement Scale (AIMS) [25]. The SAS is comprised of 10 items, each rated 0–4, subjects in the present study scored from 0–2. A score of 0 indicates normal, 1 indicates a mild level of severity, 2 indicates moderate. The AIMS is comprised of 7 individual items rated 0–4, subjects in the present study scored from 0–2. A score of 0 indicates normal, 1 indicates minimal severity, 2 indicates mild severity. Trained raters demonstrated acceptable inter-rater reliability in assessing movement disorders using a set of 15 videotaped examinations (intraclass correlation coefficient 0.93). Halifax subjects were assessed with the Extrapyramidal Syndrome Rating Scale (ESRS) [26]. The global scores are rated 0–8, subjects in the present study scored from 0–2. A score of 0 indicates normal, 1 indicates borderline and 2 indicates a definite but very mild level of severity. Inter-rater reliability for the ESRS was good (intraclass correlation coefficient 0.84).

The approaches to assessing EPS at the two sites involved the use of different, but standardized and widely accepted scales. Compared with the SAS, the ESRS contains more items assessing a variety of features of Parkinsonism and also assesses dystonia. The use of different
scales precludes pooling data between sites. However, convergent evidence obtained with different strategies also has advantages. The individual items of the SAS and AIMS, and the subscales of the ESRS assess multiple domains of extrapyramidal dysfunction. Following antipsychotic drug treatment, there is generally a significant inter-correlation between scores on individual items or subscales, and summary scores can be used for analysis. However, in never medicated subjects with schizophrenia there may be no inter-correlation between individual items or subscales [17,27], and summary scores may fail to detect subtle but definite signs of extrapyramidal dysfunction. We categorized subjects according to the presence or absence of definite signs of extrapyramidal dysfunction. Scores of 1 or more on individual items of the SAS or AIMS, and scores of 2 or more on the global subscales of the ESRS were used for this purpose.

1.3 Assessment of symptoms and treatment response

Symptom severity prior to antipsychotic drug treatment was rated using the Positive and Negative Syndrome Scale (PANSS) [28]. This is a 30 item scale, with items rated 1–7. There are three subscales, for positive and negative symptoms (7 items each), and for general psychopathology (16 items). In the Hong Kong sample, PANSS scores were available for 80/84 subjects for the acute treatment response analysis. All were treated with typical antipsychotic medications (mean 358 mg CPZ equivalents, SD 461) for a mean of 6.2 weeks (SD 4.1). PANSS scores were available for 57/84 subjects for the long term treatment response analysis (mean 254 mg CPZ equivalents, SD 252). At both time points the mean doses in subjects with and without pre-existing EPS did not differ. In the Halifax sample, acute treatment response PANSS scores were available for 28/40 subjects. All were treated with atypical antipsychotic medications (risperidone n = 17, mean dose 2.9 mg, olanzapine n = 4, mean dose 16.9 mg, quetiapine n = 7, mean dose 400 mg) for a mean of 10.3 weeks (SD 2.5). Long term treatment response data was available for 26/40 subjects. All were treated with atypical antipsychotic medications (risperidone n = 12, mean dose 2.7 mg, olanzapine n = 7, mean dose 13.6 mg, quetiapine n = 7, mean dose 393 mg) for a mean of 45.6 weeks (SD 15.8). For both cohorts, the duration of treatment for subjects with baseline EPS did not differ from the no EPS group, and medication dosages were similar between groups.

Inter-rater reliability for the PANSS was assessed at each site and found to be acceptable (intraclass correlation coefficients 0.83 Hong Kong, 0.85 Halifax).

1.4 Cognitive testing

A battery of cognitive tests was used as part of the Hong Kong protocol. Testing was carried out as soon as feasible following admission to hospital, and all subjects in the present report were tested prior to receiving any medication. For the 10 subjects with EPS at baseline, we selected 10 comparison subjects from the group with no EPS. The basis of selection was the intent to match the two groups as closely as possible for age, education, gender, and duration of untreated psychosis.

An index of overall premorbid cognitive function was assessed with the information subscale of the Wechsler Adult Intelligence Scale (WAIS-R-HK, Revised Cantonese Version, Hong Kong Psychological Society, 1989). Scaled scores were used in analysis. Memory was assessed with the logical memory and visual reproduction subscales of the Wechsler Memory Scale. In the Logical Memory Test [29] (adapted for Cantonese-speaking patients, Wong C. W., personal communication), subjects were requested to memorise a verbally presented story, then repeat the story immediately and after a 30-min delay. In the Visual Reproduction Test, subjects were instructed to memorize a visually presented design for 10 s, then to draw the design from memory immediately afterwards and following a 30-min delay. Executive functions were assessed with verbal fluency and the Modified Wisconsin Card Sorting Test (MWCST)[30] (n=8 for this test). In the verbal fluency test, subjects were required to name as many exemplars as possible from the category “animal” within 1 min. In the MWCST, subjects were asked to sort a set of 48 cards according to three sorting principles (color, shape and number). Subjects were instructed that the sorting principle would change after they made 6 consecutive correct responses. The numbers of both perseverative and total errors were monitored.

1.5 Magnetic resonance imaging

Imaging was carried out as part of the Halifax protocol. Patients were scanned as soon as possible following initial assessment. MRI scans were carried out using a Siemens Magnaton Vision 1.5T scanner. The inversion recovery sequence (TR/TE = 2000/20 ms, FOV = 200 cm, matrix 168×256) generated 18 coronal slices 4 mm thick with 1 mm gaps. The gray-white contrast ratio was 1.42, better than a spoiled gradient recall acquisition sequence (0.89). Five slices spanned the basal ganglia. A single operator outlined the caudate, putamen and globus pallidus with a previously described protocol [31,32]. Intraclass correlation coefficients for inter-rater reliability with this protocol were: caudate 0.99; putamen 0.97, globus pallidus 0.96, intracranial volume 0.99. MRI scans were available at baseline for 33 patients, 32 of whom participated in a previously reported study [31].

1.6 Statistical analysis

(i) Baseline clinical variables. Comparisons between groups with and without EPS prior to treatment used t-tests for PANSS scores. Duration of untreated psychosis was not
normally distributed, so a Mann-Whitney U-test was used.

(ii) Course of EPS during treatment. The frequencies of patients with and without EPS at baseline, and after acute and long term treatment were calculated using the definitions described above. Frequencies at each of these time points were compared with baseline frequencies using the Chi-square test.

(iii) Symptom response during treatment. Repeated measures analysis of variance was used, with main effects group (baseline EPS+, EPS−), time (baseline, outcome), and a group-by-time interaction term, with total PANSS score as the dependent measure. Due to loss of subjects with time, separate repeated measures analyses were made for acute and long term outcome time points.

(iv) Cognitive function. The adequacy of matching between the EPS+ and EPS− groups that received cognitive testing was assessed using t-tests to compare age, education, and duration of hospitalization between groups. Duration of untreated psychosis was compared with a Mann-Whitney U-test. Comparisons of cognitive performance between groups were made with the Mann-Whitney U-test.

(v) Basal ganglia structure. An analysis of covariance was used to compare volumes, with main effects group and gender, covariates intracranial volume and age, and a group-by-time interaction term [33]. Pearson correlation coefficients were calculated to investigate the relationship between age and volume of each structure, in each group (baseline EPS+, EPS−, control). A secondary analysis was carried out using a similar strategy as applied to investigate striatal structure in chronic, never mediated patients [21]. Three groups of pairs of subjects were formed (n=13/group), matched for gender and as closely as possible for age. Pairs were: EPS+ and EPS−, EPS+ and controls, EPS− and controls. Paired t-tests were carried out, as reported in the study of chronic subjects.

2 Results

2.1 EPS in subjects with no previous antipsychotic treatment

Demographic and clinical features of illness appear in Table 1. The overall prevalence of EPS was 10/84 (11.9%) in the Hong Kong sample and 17/40 (42.5%) in the Halifax sample. The predominant expression was Parkinsonism (see Table 2).

In the Hong Kong sample, EPS+ subjects had a longer duration of untreated psychosis (DUP, Mann-Whitney U-test, Z = 2.18, P = 0.03). The median DUP was also longer in the Halifax EPS+ subjects, but the difference was not statistically significant. At baseline, EPS+ subjects had higher total PANSS scores in both the Hong Kong (19.9%, t = 2.45, P = 0.016) and Halifax (14.7%, t = 2.04, P = 0.049) groups (Figure 1). The Hong Kong EPS+ group had more severe positive (t = 2.00, P = 0.049) and negative (t = 3.00, P = 0.004) symptoms than the EPS− group. Differences

| Variable                      | Hong Kong | Halifax | Controls |
|------------------------------|-----------|---------|----------|
|                              | EPS+ (mean, SD) | EPS− (mean, SD) | EPS+ (mean, SD) | EPS− (mean, SD) | n (males/females) |
| n (males/females)            | 10 (6 m/4 f) | 74 (35 m/39 f) | 17 (14 m/3 f) | 23 (17 m/6 f) | 23 (12 m/11 f) |
| Age (year)                   | 28.1 (6.4) | 32.2 (8.8) | 22.9 (4.4) | 24.0 (5.3) | 27.7 (7.2) |
| Education (year)             | 10.1 (2.0) | 10.6 (3.4) | 11.4 (1.7) | 11.8 (1.7) | 15.8 (4.6) |
| Duration of untreated psychosis (d) | 730 (median)* | 120 (median) | 309 (median, n=13) | 230 (median, n=16) |
| Ethnicity (n)                | Chinese | 10 | 74 | 16 | 19 | 22 |
|                              | Caucasian | 1 | 2 | 1 | 1 | 1 |
|                              | African-American | 1 | 1 | 1 | 1 |
|                              | First Nations | 1 | 1 | 1 | 1 |
|                              | Asian | 1 | 1 | 1 | 1 |
|                              | Other | 1 | 1 | 1 | 1 |
| Diagnosis (n)                | Schizophrenia | 8 | 37 | 16 | 21 | |
|                              | Schizoaffective | 1 | 1 | 1 | 1 |
|                              | Schizophreniform | 1 | 15 | 1 | 1 |
|                              | Brief psychosis | 8 | 8 | 8 | 8 |
|                              | Delusional disorder | 1 | 10 | 1 | 1 |
|                              | Depression with psychosis | 2 | 2 | 2 | 2 |
|                              | PNOS | 2 | 2 | 2 | 2 |

Table 1 Demographic and clinical variables for never medicated subjects with psychosis and healthy comparison subjects

a) Significant differences between EPS+ and EPS− groups are indicated *P < 0.05.
Table 2  Course of EPS with treatment a)

| Baseline status | Hong Kong       | Prevalence of EPS | Halifax        |
|-----------------|-----------------|-------------------|----------------|
|                 | Baseline | 6 weeks | 26 weeks | Baseline | 10 weeks | 46 weeks |
| Parkinsonism    | EPS+     | 10/10   | 6/9      | 2/8      | 14/17    | 1/9      | 6/11    |
| n (%)           | (100)    | (66.7)  | (25)     | (82.4)   | (11.1)   | (54.5)   |
| EPS−            | 0/74     | 25/73   | 17/50    | 0/23     | 2/14     | 1/15     |
| n (%)           | (0)      | (34.2)  | (34.0)   | (0)      | (14.3)   | (6.7)    |
| Total           | 10/84    | 31/82   | 19/58    | 14/40    | 3/23     | 7/26     |
| n (%)           | (11.9)   | (37.8)  | (32.8)   | (35.0)   | (13.0)   | (26.9)   |
| Dyskinesia      | EPS+     | 1/10    | 1/9      | 0/8      | 4/17     | 0/9      | 4/11    |
| n (%)           | (10.0)   | (11.1)  | (0)      | (23.5)   | (0)      | (36.4)   |
| EPS−            | 0/74     | 3/73    | 2/50     | 0/23     | 0/14     | 0/15     |
| n (%)           | (0)      | (4.1)   | (4.0)    | (0)      | (0)      | (0)      |
| Total           | 1/84     | 4/82    | 2/58     | 4/40     | 0/23     | 4/26     |
| n (%)           | (1.2)    | (4.9)   | (3.4)    | (10.0)   | (0)      | (15.4)   |
| Dystonia        | EPS+     | 6/17    | 2/9      | 3/11     |
| n (%)           | (35.3)   | (22.2)  | (27.3)   |
| EPS−            | 0/23    | 0/14    | 0/15     |
| n (%)           | (0)      | (0)     | (0)      |
| Total           | 6/40     | 2/23    | 3/26     |
| n (%)           | (15.0)   | (8.7)   | (11.5)   |

a) For this table, patients were split into two groups according to baseline status prior to treatment: those with (EPS+) and those without (EPS−) extrapyramidal symptoms. These group assignments were maintained, and frequencies (percents) of EPS after acute and chronic treatment were tabulated. Typical antipsychotic drugs were used in the Hong Kong sample, atypical antipsychotics in the Halifax sample.

Figure 1  Severity of symptoms at baseline for never medicated subjects with psychosis. Boxplots indicate mean (square), and 95th, 75th, 50th, 25th and 5th percentiles. Significant differences between EPS+ and EPS− groups are indicated *P < 0.05, ***P < 0.005.

between groups were similar in direction in the Halifax cohort, but were not statistically significant.

2.2 Course of EPS during treatment

Treatment was with typical antipsychotics in the Hong Kong sample. In the overall group, EPS increased in frequency following a mean of 6 weeks of treatment, and remained elevated following a mean of 26 weeks of treatment (Table 2, Figure 2). Dyskinesia developed in 2 subjects by the long term outcome point, both of whom were free of EPS at baseline. The likelihood of having EPS at either outcome time point was not related to EPS status at baseline (acute outcome Chi square = 3.25, P = 0.07; long term outcome Chi square = 0.37, P = 0.57).

Treatment was with atypical antipsychotics in the Halifax sample. In the overall group, EPS declined in frequency after a mean of 10 weeks of treatment, but increased again
after a mean of 46 weeks of treatment. Dyskinesia was present in 4/26 (15.4%) at long term follow up. Prior to antipsychotic treatment, one of these subjects had dyskinesia, three had evidence of Parkinsonism, and two had evidence of dystonia. None of the subjects who were free of EPS at baseline developed dyskinesia. At the acute treatment response time point, the frequency of EPS was not related to EPS status at baseline (Chi square = 1.17, P = 0.28). However, the likelihood of having EPS at long term outcome was related to EPS status at baseline (Chi square = 12.24, P = 0.0005). Subjects with pre-existing EPS were more likely to have EPS at long term follow up.

2.3 Baseline EPS and symptomatic treatment response

The patterns of response of symptoms are illustrated in Figure 3. In both samples, repeated measures analysis of variance indicated statistically significant differences between EPS+ and EPS− groups following acute treatment (Hong Kong F = 8.58, P = 0.005; Halifax F = 6.53, P = 0.017). Patients with EPS at baseline had higher levels of total symptoms overall. This difference continued to be statistically significant at the 26 week time point in the Hong Kong sample (F = 4.70, P = 0.03) but was not significant in the Halifax sample at 46 weeks. There were no significant interactions between group and time. Concerning the acute response of positive symptoms, in both samples there were statistically significant differences related to the presence of baseline EPS (Hong Kong F = 4.26, P = 0.043; Halifax F = 4.71, P = 0.039). There were no statistically significant differences when the long term treatment response was considered, and there were no group-by-time interactions.

Analysis of the acute treatment response of negative symptoms indicated effects related to presence of baseline EPS (Hong Kong F = 7.47, P = 0.008; Halifax F = 3.18, P = 0.09). These were persistent in the Hong Kong group only when the long term treatment response was considered (F = 6.36, P = 0.01). There were no group-by-time interactions in either sample.

2.4 Baseline EPS and cognitive function

Cognitive function was compared between demographically similar subgroups in the Hong Kong sample (Table 3). There were no differences in the index measure of premorbid global cognitive function. Two differences emerged on more specific tests, with relatively impaired performance in the EPS+ group. These were delayed recall in the visual reproduction test (approximate effect size 0.26) and increased perseverative errors in the WCST (approximate effect size 0.27). These results must however be treated with caution as the sample sizes were small and the results not corrected for multiple testing.

2.5 Baseline EPS and basal ganglia structure

At the time of MRI, 12 subjects were never medicated (4/13 with EPS, 8/20 without EPS). The mean duration of treatment for the remaining 21 subjects was 6.5 weeks (SD 3.5,
Table 3  Cognitive performance during first hospitalization in patients with EPS compared with a matched group with no EPS prior to antipsychotic drug treatment\(^a\)

| Demographics | EPS+ | EPS− |
|---------------|------|------|
| Age (year)    | 28.10| 28.50| 0.14 | 0.89 |
| Education (year) | 10.11| 10.00| 0.13 | 0.90 |
| Gender (M/F)  | 6/4  | 6/4  |      |      |
| DUP (d)       | 730  | 683  | 32.5 | 0.52 |
| Duration of hospitalization (d) | 51   | 37   | 1.01 | 0.33 |

| General cognition | EPS+ | EPS− | t-value/U-value | P-value |
|-------------------|------|------|-----------------|--------|
| Information subscale | 11.00| 11.00| 45              | 0.739  |

| Memory | EPS+ | EPS− | t-value/U-value | P-value |
|--------|------|------|-----------------|--------|
| Logical memory immediate recall (%) | 36.96| 26.09| 30              | 0.143  |
| Logical memory delayed recall (%) | 23.92| 15.22| 40              | 0.481  |
| Visual reproduction immediate recall (%) | 19.00| 20.00| 22.5            | 0.036  |
| Visual reproduction delayed recall (%) | 17.00| 19.50| 40              | 0.036  |

| Executive functions | EPS+ | EPS− | t-value/U-value | P-value |
|---------------------|------|------|-----------------|--------|
| Verbal fluency      | 19.00| 17.50| 48              | 0.912  |
| WCST perseverative error (number) | 10.00| 4.50 | 12              | 0.012  |
| WCST category       | 3.50 | 3.50 | 30.5            | 0.408  |

| a) *P<0.05. |

range 1.9–12.4), and did not differ between EPS+ and EPS− subjects. At the time of scan, 16 subjects were receiving risperidone (mean 2.9 mg), 2 received olanzapine (15 mg), 1 received quetiapine (300 mg), 1 received haloperidol (EPS−, 6 weeks, 3 mg) and 1 received a study drug (EPS−, 9 weeks, either risperidone or haloperidol).

Basal ganglia volumes are illustrated in Figure 4. Comparison of caudate volumes revealed a significant effect of group (F = 3.38, P = 0.042). Volumes were larger in the EPS+ group (4.5% vs. EPS−, 3.6% vs. healthy subjects). No statistically significant differences were observed for putamen or globus pallidus volumes. The correlations between volume and age appear in Table 4. Caudate volume was negatively correlated with age in the EPS+ group only (P = 0.017). Putamen volume was negatively correlated with age in the healthy comparison group (P = 0.0006), and globus pallidus volume was also negatively correlated with age in this group (P = 0.012). Correlations between putamen or globus pallidus volumes and age were not statistically significant in either EPS group.

In subjects who participated in the MRI scan, 14/20 EPS− had a lifetime history of exposure to marijuana, versus 4/13 EPS+ (Chi-square = 4.89, P = 0.027). Overall, when marijuana exposure was substituted for EPS in the model, subjects with marijuana exposure did not show a difference in caudate, putamen or globus pallidus volumes compared with those who had no exposure. In subjects who participated in the MRI scan, 1/20 EPS− had a lifetime history of previous exposure to stimulant drugs, versus 1/13 EPS+. When these two subjects were excluded from the overall analysis, the statistical significance of the results was unchanged.

The paired analysis was carried out to replicate the strategy used to investigate volume differences between chronic, never medicated patients with and without movement disorders [21]. The results were similar to the analysis above for the caudate, and additional findings emerged for the putamen and the globus pallidus (Figure 5). For the caudate, volumes of the EPS+ group were 6.8% larger than the EPS− group (paired t = 2.22, P = 0.047). For the putamen, the EPS− group volumes were 4.4% smaller than healthy comparison subjects (paired t = 2.39, P = 0.033). For the globus
Subjects [18,20,22]. In both cohorts, higher total symptom dyskinesia may overlap significantly in never medicated samples, and Parkinsonism and tardive dyskinesia may represent progression of the movement disorder process, or development of tolerance to a suppressive effect of antipsychotic drugs. A similar observation was made in the Hillside Hospital study [16,36]. In the Hong Kong cohort, baseline EPS appeared to decline over the first 6 months of typical antipsychotic treatment. In other studies, similar observations were reported for neurological soft signs [37,38].

Subjects in the Halifax cohort were treated with atypical antipsychotics. Subjects with EPS prior to treatment had reduced EPS after 10 weeks of treatment, similar to a report of 2–14 weeks of treatment with risperidone in subjects with baseline EPS [39]. A longer term study of subjects with baseline EPS indicated no change in overall severity after 6 months of treatment, largely with atypical antipsychotics [34]. In the present study, later re-emergence of EPS in predisposed subjects after a mean of 46 weeks may represent progression of the movement disorder process, or development of tolerance to a suppressive effect of medications. As well, the initial dopamine receptor occupancy may eventually result in receptor up regulation which differs with different types of antipsychotic drugs [40].

The treatment response of psychopathology was less complete in EPS+ subjects in both cohorts in the present study. These results are similar to two other reports, where patients with baseline EPS present were less likely to respond to treatment, and when they did respond, this took longer [16,18]. However, these observations are in contrast to another study, where Parkinsonism at baseline appeared to predict extent of response of positive symptoms [34].

We observed poorer executive function in subjects with baseline EPS. There are few other studies of cognitive function in subjects with EPS prior to treatment. Older, never medicated subjects with dyskinesia appeared to have similar memory dysfunction on the Wechsler Memory Scale as those with no dyskinesia [23]. Global cognitive impairment

### Table 4 Correlations between age and volumes of basal ganglia structures

|                | Caudate          | Putamen         | Globus pallidus |
|----------------|------------------|-----------------|-----------------|
| n              |                  |                 |                 |
| EPS+           | 13               | -0.64*          | -0.88, -0.14    |
|                | 20               | 0.07            | -0.64*          |
| Controls       | 23               | -0.18           | -0.55, 0.25     |

*These were statistically significant *P<0.05 for caudate in the EPS+ group, and for putamen and globus pallidus in the control group.

The present results linking pre-treatment EPS to symptom severity in first episode psychosis, to cognitive dysfunction, and to abnormalities of brain structure all support the proposal that similar observations in family members of patients with psychosis represent an endophenotype of the illness [5,6]. Findings concerning the course of extrapyramidal symptoms and the relationships to treatment response indicate the possible importance of thorough assessment for EPS at the entry into early psychosis treatment programs.

The prevalence of EPS was within the range reported in early psychosis [9–18,34,35]. We used the presence of any EPS to form a subgroup for analysis. While subgrouping according to type of EPS would have been preferable, the sample size was not large enough, and Parkinsonism and dyskinesia may overlap significantly in never medicated subjects [18,20,22]. In both cohorts, higher total symptom severity scores were associated with pre-treatment EPS. In the Hong Kong cohort, both positive and negative symptom scores were also greater in the EPS+ group. These relationships are similar to those reported in the literature for total 16–18,34]. In chronic never-medicated subjects, these relationships were not observed [20]. One of our cohorts demonstrated a relationship between severity of EPS and duration of untreated psychosis, not seen elsewhere [16,17].

Risk of treatment-related Parkinsonism was unrelated to pre-existing EPS in the Hong Kong sample, where the acute treatment dose of typical antipsychotics was approximately 360 mg CPZ equivalents/d. In contrast, pre-treatment EPS increased the risk for Parkinsonism in the acute phase of treatment with 20–40 mg/d of fluphenazine or haloperidol (approximately 1000–2000 CPZ equivalents) [16]. We did not find an association between pre-treatment EPS and risk for developing tardive dyskinesia related to typical antipsychotic drugs. A similar observation was made in the Hillside Hospital study [16,36]. In the Hong Kong cohort, baseline EPS appeared to decline over the first 6 months of typical antipsychotic treatment. In other studies, similar observations were reported for neurological soft signs [37,38].

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Subjects in the Halifax cohort were treated with atypical antipsychotics. Subjects with EPS prior to treatment had reduced EPS after 10 weeks of treatment, similar to a report of 2–14 weeks of treatment with risperidone in subjects with baseline EPS [39]. A longer term study of subjects with baseline EPS indicated no change in overall severity after 6 months of treatment, largely with atypical antipsychotic drugs [34]. In the present study, later re-emergence of EPS in predisposed subjects after a mean of 46 weeks may represent progression of the movement disorder process, or development of tolerance to a suppressive effect of medications. As well, the initial dopamine receptor occupancy may eventually result in receptor up regulation which differs with different types of antipsychotic drugs [40].

The treatment response of psychopathology was less complete in EPS+ subjects in both cohorts in the present study. These results are similar to two other reports, where patients with baseline EPS present were less likely to respond to treatment, and when they did respond, this took longer [16,18]. However, these observations are in contrast to another study, where Parkinsonism at baseline appeared to predict extent of response of positive symptoms [34].

We observed poorer executive function in subjects with baseline EPS. There are few other studies of cognitive function in subjects with EPS prior to treatment. Older, never medicated subjects with dyskinesia appeared to have similar memory dysfunction on the Wechsler Memory Scale as those with no dyskinesia [23]. Global cognitive impairment

3 Discussion

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In the chronic subjects, the left lentiform nucleus was also disrupted correlations between age and basal of larger caudate nuclei and possibly larger globus pallidus, parents with schizophrenia demonstrated smaller caudate volumes [54–56]. One report of adolescent offspring of healthy subjects [31,33,47–53], and others report smaller episode psychosis show no differences in volume from pre-treatment EPS. Some studies of the basal ganglia in first ing agents including cocaine and methamphetamine also though several studies may provide clues. Dopamine deplet- atum [45], an effect which is further modulated by COMT genotype [46].

This is the first MRI study separating groups of first episo- de patients according to the presence or absence of pre-treatment EPS. Some studies of the basal ganglia in first episode psychosis show no differences in volume from healthy subjects [31,33,47–53], and others report smaller volumes [54–56]. One report of adolescent offspring of parents with schizophrenia demonstrated smaller caudate volumes in the at-risk probands [57]. The present findings of larger caudate nuclei and possibly larger globus pallidus, as well as disrupted correlations between age and basal ganglia volumes associated with pre-treatment EPS are consist- ent with a study of chronic, never medicated subjects [21]. In the chronic subjects, the left lentiform nucleus was enlarged in those with EPS compared with controls [21]. The predominant form of EPS was dyskinesia, however elevated scores for Parkinsonism were also observed. Rela- tively larger caudate nuclei were also reported in subjects with chromosome 22q deletion syndrome, a significant ge- netic risk factor for schizophrenia [58–60].

The mechanism of volume enlargement is unknown, al- though several studies may provide clues. Dopamine deplet- ing agents including cocaine and methamphetamine also appear to be associated with basal ganglia enlargement [61–64]. Typical antipsychotic drug treatment enlarges basal ganglia volume [55,65,66], and these changes can be reversed by switching to atypical antipsychotics including clozapine and olanzapine [32,55,65–68]. Striatal dopamine synthesis in never medicated subjects with schizophrenia was generally reported to be increased [69–71]. However, individual, never medicated subjects with catatonia in sev- eral studies were reported to have lower dopamine synthesis than healthy subjects [69,70]. Treatment with haloperidol reduced dopamine synthesis in striatum and in thalamus, with associated increase in EPS [72]. Another study of medicated subjects indicated dorsal striatal regions may show relatively normal dopamine synthesis, while ventral regions show increased synthesis [45]. In this study, although regional volumes were not significantly different from healthy comparison subjects, the mean dorsal striatal volumes were larger than controls, while mean ventral striatal volumes were smaller. The possibility of regional varia- tion in dopamine dysfunction within the striatum in schizo- phrenia is not without precedent. In Parkinson’s disease, dopamine depletion is more pronounced in the putamen and posterior regions [73], while in methamphetamine abuse de- pletion is more notable in the caudate and dorsal regions [62].

The present study has a number of limitations. The as- certainment procedures were different at the two sites, the Hong Kong sample was limited to patients requiring inpa- tient treatment, while the Halifax sample included patients entering care as in- or out-patients. The Hong Kong sample was nearly equally split between males and females, while the Halifax sample was only one-quarter female. The origin of the gender differences is unclear, other studies from the Hong Kong EASY program also are gender balanced [74], and studies from Halifax as well as other Vancouver, Can- ada are male predominant [75,76]. Different clinical instru- ments were used between sites for assessment of several key clinical features of illness. This could contribute to discrepant findings between sites, however, has the advantage of strengthening the confidence in convergent evidence be- tween sites. Specifically, the two programs used different assessment scales for movement disorders. These may have somewhat different properties, the Simpson-Angus scale for example is highly weighted towards rigidity as a defining characteristic of Parkinsonism, and lacks an item for brady- kinesia. However, the similar associations between baseline EPS, clinical symptoms and response in both cohorts sug- gest these findings are robust. Similar to others [16], we used relatively lenient criteria to define the presence of EPS. Negative symptoms of schizophrenia are often elevated in groups with EPS, and there is likely some overlap in defini- tions. However, negative symptoms were also reported to be correlated with dystonia and dyskinesia scores [17], which are unlikely to be a consequence of observational confusion. Finally, the cognitive testing results in particular are from a small number of subjects. Although a carefully matched group was available for comparison, these results remain preliminary.

The present findings could have implications for under- standing the mechanism of illness in schizophrenia. Dyski- nesia in chronic, never medicated patients was associated with an increased frequency of dyskinesia but not Parkin- sonism in family members [77]. However, even in chronic, never treated patients, movement disorders wax and wane, and may not be a stable trait phenomena [22]. Schizotypal subjects may provide a “trait” reference point, and interestingly appear to have smaller caudate and putamen volumes [78,79]. In contrast, in most studies of first episode subjects, striatal volumes do not show statistically significant differences from healthy subjects [31,33,47–53]. Extrapyramidal symptoms and signs, as well as related cognitive dysfunc- tion may be dimensional rather than typological features of schizophrenia. Severity may be state-dependent. The rela- tionships between EPS and brain structures are likely to be moderated by treatment, and possibly by phase of illness as well. First episode studies, and investigation of subjects treated with medications that do not affect EPS, cognition or brain structure will allow better understanding of these ill- ness domains in schizophrenia.
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