Structural Brain MRI Trait Polygenic Score Prediction of Cognitive Abilities

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Structural brain magnetic resonance imaging (MRI) traits share part of their genetic variance with cognitive traits. Here, we use genetic association results from large meta-analytic studies of genome-wide association (GWA) for brain infarcts (BI), white matter hyperintensities, intracranial, hippocampal, and total brain volumes to estimate polygenic scores for these traits in three Scottish samples: Generation Scotland: Scottish Family Health Study (GS:SFHS), and the Lothian Birth Cohorts of 1936 (LBC1936) and 1921 (LBC1921). Five brain MRI trait polygenic scores were then used to: (1) predict corresponding MRI traits in the LBC1936 (numbers ranged 573 to 630 across traits), and (2) predict cognitive traits in all three cohorts (in 8,115–8,250 persons). In the LBC1936, all MRI phenotypic traits were correlated with at least one cognitive measure, and polygenic prediction of MRI traits was observed for intracranial volume. Meta-analysis of the correlations between MRI polygenic scores and cognitive traits revealed a significant negative correlation (maximal $r = 0.08$) between the HV polygenic score and measures of global cognitive ability collected in childhood and in old age in the Lothian Birth Cohorts. The lack of association to a related general cognitive measure when including the GS:SFHS points to either type 1 error or the importance of using prediction samples that closely match the demographics of the GWA samples from which prediction...
Structural brain MRI traits and cognitive abilities are heritable, with over 50% of the variance for some MRI traits — for example, frontal lobe volumes and white matter hyperintensities — being due to genes (Deary et al., 2009; Peper et al., 2007). Some of these MRI traits have been shown to share genetic variance with cognitive measures (Betjemann et al., 2010; Bohlken et al., 2014; Posthuma et al., 2002). Here, we test whether the additive effect of common DNA single nucleotide polymorphisms (SNPs) influencing cerebral white matter hyperintensities burden (WMH), brain infarcts (BI), hippocampal (HV), total brain (TBV) and intracranial (ICV) volumes predict variance in measures of cognitive ability. These MRI polygenic scores will be based on the results of four GWA studies (Bis et al., 2012; Debette et al., 2010; Fornage et al., 2011; Ikram et al., 2012), and estimated in three Scottish cohorts who have been measured on processing speed, memory, verbal, and executive function. First, we will establish whether the brain MRI polygenic scores predict their respective MRI trait in one of the cohorts who have MRI data. Where this is confirmed, we expect that common SNPs influencing these MRI traits will explain variance in the cognitive traits.

Various brain MRI structural traits are associated with cognitive ability (Andreasen et al., 1993; Haier et al., 2004). The most investigated of these is TBV, which correlates 0.33 with intelligence, as estimated from a meta-analysis of 37 samples (n = 1,530; McDaniel, 2005). Twin studies have supported complete genetic mediation of this relationship in adults (Posthuma et al., 2002); and in children, genetic overlap has been shown between measures of TBV, neocortex, white matter, and pre-frontal cortex with a range of cognitive indices (IQ, reading ability, processing speed; Betjemann et al., 2010). ICV, which might be considered as a pre-morbid/maximal brain size measure, has been associated with vocabulary performance (Schottenbauer et al., 2007), and with semantic memory, executive function, and spatial ability when adjusting for current brain pathology in older people (Farias et al., 2012). HV has largely been investigated in relation to memory abilities. A meta-analysis of 33 studies reporting correlations between HV and memory performance showed a negative correlation of 0.25 for children and young adult samples, and a positive correlation (0.10) in older samples (Van Petten, 2004). Heterogeneity within older sample estimates indicated a variable association dependent on age-related changes, possibly influenced more by environmental factors, which have a greater effect on HV in old age than do genes (Sullivan et al., 2001).

Other brain MRI traits have shown significant associations with particular cognitive domains or in specific demographic groups. WMH, for example, are mainly associated with impaired executive functioning, particularly in aging populations where WMH are more prevalent (Farias et al., 2012; Gunning-Dixon & Raz, 2000; Hedden et al., 2012). A twin study of older men showed that 70–100% of the correlation between WMH and cognitive traits was due to common genes (Carmelli et al., 2002). BIs are also related to cognitive dysfunction and decline in the elderly, with rates being increased even in persons with covert BI in the absence of clinical stroke events (Vermeer et al., 2003). The genetic underpinning of this relationship is unknown.

The genetic covariance between brain MRI and cognitive traits provides the rationale for our investigation, which aims to establish whether the variability in cognition can be partly explained by structural brain differences. No common genes of large effect (e.g., >5% variance) have been reported for brain MRI traits. Therefore, we created brain MRI polygenic scores based on the summative influence of SNPs with differing levels of effect size (i.e., from significant to non-significant effects) from recent GWA meta-analysis studies (Bis et al., 2012; Debette et al., 2010; Fornage et al., 2011; Ikram et al., 2012). We tested whether these polygenic scores are predictive of (1) their respective MRI trait, and (2) of cognitive variation.

Material and Methods
Cohorts
Brain MRI polygenic profile scores were calculated in three independent Scottish cohorts: GS:SFHS, the LBC1936, and the LBC1921. GS:SFHS is a large population family-based study of around 24,000 Scottish participants sampled between the years 2006 and 2011 (www.generationscotland.org/); 10,000 participants were selected for genome-wide analysis based on: Caucasian ethnicity, being born in the United Kingdom and full phenotype data (Kerr et al., 2013). In the current analysis, only unrelated subjects were included, leaving an analysis sample of 6,814. The mean age of the sample was 55.5 years (SD = 11.4) at testing (59% women). The LBC samples comprised relatively healthy individuals born in 1921 or 1936 in the Edinburgh area, most of whom had completed the Moray House Test No. 12 (MHT) assessment of general intelligence in the Scottish Mental Surveys of 1932 or 1947 at

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a mean age of 11 years (Deary et al., 2012). The LBC1936 (n = 1,091; 49.8% women) were tested on the MHT and other cognitive measures in adulthood at a mean age of 69.5 years (SD = 0.8). At age 73 years, a subset of these individuals (n = 724) underwent structural MRI. The LBC1921 (n = 550; 57.4% women) completed the MHT plus additional cognitive tests at a mean age of 79.1 years (SD = 0.6) and later at 83.3 years (SD = 0.54). Following informed consent, venesected whole blood was collected for DNA extraction for the LBC samples, with both saliva and blood being used for DNA extraction in the GS:SFHS. Ethical approval for the LBCs was obtained from Scotland’s Multicentre Research Ethics Committee and local research ethics committee. GS:SFHS ethical approval was granted by the NHS Tayside Committee on Medical Research Ethics (REC Reference Number: 05/S1401/89). Research Tissue Bank status was approved by the Tayside Committee on Medical Research Ethics (REC Reference Number: 10/S1402/20), enabling generic ethical approval for medical research purposes.

MRI Measures in LBC1936
Structural T2-, T2*-, FLAIR- and T1-weight brain MRI data were collected using a GE Signa 1.5 T HDX clinical scanner. Bs were coded for size and location based on vascular territory and typical signal characteristics by consultant neuroradiologists using a validated stroke lesion rating scale (Wardlaw & Sellar, 1994; Wardlaw et al., 2011), which differentiates infarcts into cortical, lacunar, borderzone, and brainstem/cerebellar. Lacunar infarcts were coded as being cavitated or not (Wardlaw et al., 2013).

BI of any size and location were present in 95 individuals and absent in 537. WMH measured in the white matter and subcortical grey matter, including cerebellum and brainstem, was quantified semi-automatically with MCMxxxVI (Hernandez Mdel et al., 2010). Images were inspected and false positive and negative lesions manually corrected (http://www.bric.ed.ac.uk/research/imagereanalysis.html).

Focal stroke lesions were masked manually to distinguish them from other structures. The dependent measure was the natural logarithm (WMH burden in mL +1). ICV includes the contents within the inner skull table, including venous sinuses, and has its inferior limit in the axial slice just superior to the tip of the odontoid peg at the foramen magnum and superior to the inferior limits of the cerebellar tonsils. The ICV was obtained semi-automatically using the T2*W sequence. The first approximation of the ICV was obtained automatically using the Object Extraction Tool in Analyze 9.0. Then, the cervical spinal cord inferior to the inferior boundary was removed manually, along with the pituitary gland (in cases where this latter structure was included). HV was obtained after an automatic segmentation of left and right hippocampi using FSL tools (www.fmrib.ox.ac.uk/fsl) and an ageing template. The resulting automatically segmented masks were visually assessed for accuracy and manually edited using Analyze 9.0 (Mayo Clinic, AnalyzeDirect, Inc. Mayo Clinic) if required. Mean of left and right HV was used. TBV (mm$^3$) was defined by the volume of the cerebrosinal fluid, venous sinuses, and meninges subtracted from the ICV. To correct for variation in head size between individuals, HV and TBV were expressed as percentages of ICV.

Cognitive Measures Collected in All Cohorts
In GS:SFHS, four cognitive ability tests were administered: the Wechsler Digit Symbol Substitution Test (DS), Wechsler Memory Scale Logical Memory Test (sum of immediate and delayed recall of one paragraph) (LM), the phonemic Verbal Fluency Test using the letters C, F, and L, each for one minute (VF), and the Mill Hill Vocabulary Scale combining junior and senior synonyms (MHV; Smith et al., 2006). In the LBC samples, DS, LM, and VF tests were administered, but instead of the MHV, the National Adult Reading Test (NART; Nelson, 1982) was used to index vocabulary ability. For the LBC1921, DS was only measured at age 83 years, where the sample size was reduced (n = 302). A composite score of these four measures (or three age 79 years measures for the LBC1921) was formed by deriving regression-based factor scores from an unrotated principal components analysis that explained 45–55.3% of variance across cohorts. In addition to these four overlapping tests across the three cohorts, LBC samples had overlapping MHT scores from childhood (MHT11) and old-age (MHT).

Genotyping
Genotyping in the GS:SFHS and LBC samples was performed at the Wellcome Trust Clinical Research Facility Genetics Core, Edinburgh (www.wtcrf.ed.ac.uk). GS:SFHS samples were genotyped on the Illumina HumanOmniExpressExome-8 v1.0 DNA Analysis BeadChip using Infinium chemistry (Marioni et al., 2014). In the LBC samples, Illumina Human610-QuadV1 Chip whole genome genotyping was available. Genotype quality control procedures are described elsewhere (Davies et al., 2011), but briefly, necessary exclusion were made for gender discrepancies, individual relatedness, and non-Caucasian descent. Good quality genotyping information was available for 509 (LBC1921) and 1005 (LBC1936) Caucasian individuals.

Statistical Analysis
Five sets of brain MRI polygenic scores — BI, WMH, ICV, TBV, and HV — were estimated using SNP association procedures are described elsewhere (Davies et al., 2011), but briefly, necessary exclusion were made for gender discrepancies, individual relatedness, and non-Caucasian descent. Good quality genotyping information was available for 509 (LBC1921) and 1005 (LBC1936) Caucasian individuals.
Structural Brain MRI Trait Polygenic Score Prediction

For the cognitive measures, similar regression models were tested, but with age at MRI scanning replaced by age at cognitive test. Standardized betas from the regression models for the cognitive traits were meta-analyzed under a random effects model in R (MAc package; R Development Core Team, 2008) giving an overall effect size and standard error. Given the intercorrelations among the four cognitive tests and among the five MRI polygenic traits, we made an alpha-level adjustment based on a matrix spectral decomposition (Nyholt, 2005) of these traits (g was not included because it is a composite measure of the cognitive tests, and we chose one polygenic score \((p < 1\) inclusion threshold) to avoid dependency among polygenic scores at differing SNP p-value inclusion levels). Using the largest cohort, GS:SFHS, we identified 8.86 effective traits to give an adjusted alpha level of 0.006. Heterogeneity between sample estimates was tested via Cochran’s Q statistic.

Results

Polygenic Prediction of MRI Traits in the LBC1936

The sample size varied between 573 (WMH) and 629 (BI). ICV was the only variable to show significant associations at the corrected alpha level \((p < .01\); correlations ranged from 0.08–0.10 across all p value polygenic inclusion criteria (Table S4). The HV polygenic score was most strongly correlated with HV at the polygenic \(p < .05\) inclusion \((r = 0.08, p = .04)\) and polygenic \(p < 1\) inclusion \((r = 0.07, p = .05)\). Polygenic scores for TBV at the polygenic \(p < .01\) inclusion was correlated 0.08 with TBV \((p = .02)\). For WMH, correlations of 0.08 and 0.09 \((p < .05)\) were observed at \(p < .05\), \(p < .50\), and \(p < 1\) polygenic inclusion thresholds. All BI polygenic score correlations showed correlational p values greater than 0.05 with BI. Exclusion of stroke cases did not alter the polygenic score effects; thus, subsequent results are reported for the larger sample to reduce the standard error of the estimates.

Correlation Between MRI and Cognitive Traits in LBC1936

Phenotypic correlations between MRI traits and the main cognitive traits of interest in the LBC1936 \((n = 629)\) are shown in Table S3. With the exception of HV, all brain MRI traits were significantly correlated with at least one cognitive trait. BI correlated significantly (negatively) with all cognitive traits \((n = 625–629)\) and TBV correlated significantly (positively) with all traits except NART \((n = 619–623)\).

Polygenic Prediction of Cognitive Traits in All Cohorts

Meta-analysis results of the correlations between the brain MRI polygenic scores (at differing polygenic p-value inclusion intervals) and cognitive measures are shown in Table 1. The only significant correlations (at the corrected alpha level) to demonstrate consistency across differing polygenic scores at each SNP inclusion criterion \((p < .01, p < .05, p < .1, p < .5, p < 1)\). Because the polygenic scores at different SNP inclusion are non-independent, we made a Bonferroni correction to our alpha level of 0.05 for the five polygenic MRI traits, which gave an adjusted significance level of 0.01.
$p$-value threshold scores for HV with MHT11 and MHT in old age (at polygenic $p < .50$ and 1 thresholds). For MHT11, the correlational $p$ value was .003 and for MHT, it was .003. A negative correlation was observed such that a smaller HV was related to better MHT scores. Forest plots for these variables (only measured in the LBC samples) and for a comparable measure (general cognitive ability) in all three cohorts are shown in Figure S1. For TBV, heterogeneity was found between cohort estimates for DS, MHT11, and MHT.

**Discussion**

Our study showed that MRI ICV polygenic scores derived from GWA results on around 10,000 individuals (CHARGE) were predictive of variance in ICV in 624 subjects aged 72 years. On a phenotypic level, BI and WMH were negatively correlated with cognitive measures in this cohort, whereas the cranial and TBV measures were positively correlated. A meta-analysis of this and another elderly cohort showed HV polygenic scores were negatively correlated with the same general cognitive ability test measured in childhood and old age, explaining up to 1% of cognitive core variance. No other brain MRI polygenic scores were significantly associated with any other cognitive traits in the meta-analyses, including all three Scottish cohorts.

This is the first study to test whether the SNP effects reported in current GWA studies of brain MRI traits are predictive of variance in these traits in an independent sample. At a significance level corrected for multiple testing, we confirmed this for ICV and for other traits (WMH, TBV, and HV) at an unadjusted alpha level of 0.05. The amount of variance explained by these polygenic effects, although small, is consistent with other polygenic prediction studies of psychiatric and disease traits and is argued to increase with increases in size of the GWA samples on which prediction is based (Dudbridge, 2013). The lack of polygenic prediction for BI is likely due to the small number of

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**TABLE 1**

<table>
<thead>
<tr>
<th>$p$-value</th>
<th>Digit symbol (N = 8,020)</th>
<th>Verbal fluency (N = 8,250)</th>
<th>Logical memory (N = 8,249)</th>
<th>Vocabulary (N = 8,212)</th>
<th>General factor (N = 8,115)</th>
<th>MHT11 (N = 1,411)</th>
<th>MHT (N = 1,498)</th>
</tr>
</thead>
<tbody>
<tr>
<td>BI</td>
<td>.01 (0.01)</td>
<td>(0.01)</td>
<td>(0.01)</td>
<td>(0.01)</td>
<td>-0.06 (0.06)</td>
<td>-0.11 (0.11)</td>
<td>-0.04 (0.03)</td>
</tr>
<tr>
<td>.05</td>
<td>(0.01)</td>
<td>(0.01)</td>
<td>(0.01)</td>
<td>-0.01 (0.01)</td>
<td>0.01 (0.01)</td>
<td>-0.02 (0.03)</td>
<td>0.03 (0.03)</td>
</tr>
<tr>
<td>.10</td>
<td>(0.01)</td>
<td>(0.01)</td>
<td>(0.01)</td>
<td>-0.02 (0.01)</td>
<td>(0.02)</td>
<td>-0.05 (0.05)</td>
<td>0.00 (0.03)</td>
</tr>
<tr>
<td>.50</td>
<td>(0.01)</td>
<td>(0.01)</td>
<td>(0.01)</td>
<td>-0.03 (0.01)</td>
<td>-0.02 (0.01)</td>
<td>-0.04 (0.03)</td>
<td>-0.02 (0.03)</td>
</tr>
<tr>
<td>1</td>
<td>(0.01)</td>
<td>(0.01)</td>
<td>(0.01)</td>
<td>-0.02 (0.01)</td>
<td>-0.01 (0.02)</td>
<td>-0.04 (0.03)</td>
<td>0.00 (0.03)</td>
</tr>
</tbody>
</table>

**WMH**

| .01       | (0.01)                    | (0.01)                     | (0.01)                      | (0.01)                 | (0.01)                   | 0.01 (0.03)     | 0.00 (0.03)    |
| .05       | (0.01)                    | (0.01)                     | (0.01)                      | (0.01)                 | (0.01)                   | 0.01 (0.04)     | 0.00 (0.05)    |
| .10       | (0.01)                    | (0.01)                     | (0.01)                      | (0.01)                 | (0.01)                   | -0.01 (0.01)    | -0.05 (0.05)   |
| .50       | (0.01)                    | (0.01)                     | (0.01)                      | (0.01)                 | (0.01)                   | -0.02 (0.01)    | -0.03 (0.03)   |
| 1         | (0.01)                    | (0.01)                     | (0.01)                      | (0.01)                 | (0.01)                   | -0.02 (0.02)    | -0.02 (0.03)   |

**ICV**

| .01       | (0.01)                    | (0.01)                     | (0.01)                      | (0.01)                 | -0.01 (0.01)             | 0.02 (0.03)     | -0.03 (0.03)   |
| .05       | (0.01)                    | (0.01)                     | (0.01)                      | (0.01)                 | (0.01)                   | 0.03 (0.03)     | -0.03 (0.03)   |
| .10       | (0.01)                    | (0.01)                     | (0.01)                      | (0.01)                 | (0.01)                   | -0.01 (0.01)    | -0.05 (0.03)   |
| .50       | (0.01)                    | (0.01)                     | (0.01)                      | (0.01)                 | (0.01)                   | -0.02 (0.02)    | -0.02 (0.03)   |
| 1         | (0.01)                    | (0.01)                     | (0.01)                      | (0.01)                 | (0.01)                   | -0.01 (0.01)    | -0.02 (0.02)   |

**HV**

| .01       | (0.02)                    | (0.02)                     | (0.02)                      | (0.02)                 | -0.01 (0.04)             | -0.01 (0.02)    | -0.04 (0.03)   |
| .05       | (0.01)                    | (0.01)                     | (0.01)                      | (0.01)                 | (0.01)                   | -0.01 (0.01)    | -0.04 (0.03)   |
| .10       | (0.01)                    | (0.01)                     | (0.01)                      | (0.01)                 | (0.01)                   | -0.01 (0.01)    | -0.05 (0.03)   |
| .50       | (0.01)                    | (0.01)                     | (0.01)                      | (0.01)                 | (0.01)                   | -0.02 (0.02)    | -0.02 (0.03)   |
| 1         | (0.01)                    | (0.01)                     | (0.01)                      | (0.01)                 | (0.01)                   | -0.01 (0.01)    | -0.02 (0.02)   |

**TBV**

| .01       | (0.03)                    | (0.03)                     | (0.03)                      | (0.03)                 | (0.03)                   | (0.03)         | (0.03)         |
| .05       | (0.03)                    | (0.03)                     | (0.03)                      | (0.03)                 | (0.03)                   | (0.03)         | (0.03)         |
| .10       | (0.03)                    | (0.03)                     | (0.03)                      | (0.03)                 | (0.03)                   | (0.03)         | (0.03)         |
| .50       | (0.03)                    | (0.03)                     | (0.03)                      | (0.03)                 | (0.03)                   | (0.03)         | (0.03)         |
| 1         | (0.03)                    | (0.03)                     | (0.03)                      | (0.03)                 | (0.03)                   | (0.03)         | (0.03)         |

Note: Bolded rows indicate polygenic scores that significantly predicted the accompanying MRI phenotype and for which we would expect significant correlations.

$p < .05, \quad ^{*}p < .01$

Sample heterogeneity ($p < .05$):

BI: MHT11 (p = .01 inclusion), LBC1936 (r = -.01), LBC1921 (r = -.13*), Verbal fluency (p < .1 inclusion) GS (r = 0), LBC1936 (r = .07*), LBC1921 (r = 0.2).

HV: Digit symbol (p < .1 inclusion), GS (r = .01), LBC1936 (r = -.09*), LBC1921 (r = -0.01)

TBV: Digit symbol (p < .01, .1, .5, 1), GS (r = 0.01, 0.01, 0.01, 0.01), LBC1936 (r = 0.11, 0.09*, .10*, .09*), LBC1921 (r = 0.04, 0.03, 0.05, 0.05), MHT11 (p < .01, .05, .1, .5, 1), LBC1936 (r = 0.07*, .05, 0.06, 0.07*, 0.07*), LBC1921 (r = -.10*, -.11*, -0.11, -.08, -.08), MHT (p < .05, .1, .5, .1), LBC1936 (r = 0.02, 0.03, 0.06, 0.06), LBC1921 (r = -.11*, -.11*, -.09*, -.10*).
individuals in our sample with BI (14.8%); as a dichotomous variable this analysis was less powered than those of continuous traits.

HV polygenic scores showed the strongest positive correlations with HV at polygenic $p < .5$ and $p < 1$ inclusion thresholds (although not significant at a corrected alpha level). It was at these thresholds that we observed a significant (negative) correlation (at a corrected alpha level) between polygenic variation in HV and phenotypic variation in MHT despite a lack of association between phenotypic variation in HV and MHT. This is an interesting finding, given that in young adulthood HV shows a negative phenotypic correlation at least with measures of memory, an aspect of general cognitive ability. Incomplete synaptic pruning during childhood and adolescence has been offered as an explanation for the negative association between HV and cognition in earlier life (Foster et al., 1999); and if genes influence pruning then it might be that this variation is driving the negative correlation between HV polygenic scores and general cognitive ability in our sample.

Alternatively, the HV polygenic score derived in our study might not be a valid measure of variation in HV because it did not significantly predict HV; therefore, any correlation with cognition could be a false positive finding. That a similar measure of general cognitive ability was not found to be associated with HV polygenic scores in the meta-analysis of all cohorts further supports this finding representing type 1 error. If it is important for the independent prediction samples to closely match the samples used in the GWA study on which polygenic scores are based, then the LBC samples more closely matched the brain MRI GWAS samples in age (being elderly), whereas GS:SFHS was predominantly comprised of individuals under the age of 60 years (62%). The polygenic scores would therefore represent genetic effects that are important in old age, so it follows that prediction is going to be more reliable in older adults. However, the observation in the LBC samples that HV polygenic scores did not predict vocabulary or BI (14.8%); as a dichotomous variable this analysis was less powered than those of continuous traits.

In conclusion, polygenic effects on MRI ICV, determined in a relatively small GWA study, were predictive of phenotypic trait variation in ICV in an independent sample. The lack of association between ICV polygenic scores and cognitive ability in the larger meta-analysis sample might suggest that other types of genetic variants (e.g., rare, structural) explain their genetic covariance. Larger GWA studies of WMH, TBV, HV, and BI will likely improve the polygenic prediction of these traits in independent samples. Polygenic scores based on these larger studies should then be investigated in relation to cognition. Improvements in the harmonization of imaging measures across studies will also enable GWA results for other brain MRI measures such as laterality to be included.

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Conflict of Interest
None.

Ethical Standards
The authors assert that all procedures contributing to this work comply with the ethical standards of the relevant national and institutional committees on human experimentation and with the Helsinki Declaration of 1975, as revised in 2008.

Supplementary Material
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