‘I'm always up against a brick wall with them’ : parents' experiences of accessing support for their child with a newly recognised developmental disorder

McCarthy, RE, Blackburn, C, Mukherjee, RAS, Fleming, K, Allely, CS, Kirby, L and Cook, PA

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‘I’m always up against a brick wall with them’: parents’ experiences of accessing support for their child with a newly recognised developmental disorder

Robyn McCarthy, Carolyn Blackburn, Raja Mukherjee, Kate Fleming, Clare Allely, Lauren Kirby and Penny A. Cook

Three of the most prevalent developmental disorders (DDs) are autism spectrum disorder (ASD), attention-deficit/hyperactivity disorder (ADHD) and fetal alcohol spectrum disorder (FASD). As part of a study screening for DDs in Greater Manchester, UK, a unique opportunity was taken to explore parents’ experiences of receiving reports about their child’s previously unrecognised DD. Six parents out of a possible 16 took part in interviews, which were analysed thematically. Findings drawn from parental responses revealed a number of barriers to accessing support for their child’s additional needs, including perceived resistance from schools, particularly for quiet, well-behaved girls, and difficulty in accessing assessment or referrals. There needs to be greater awareness of additional needs in children without externalising behaviours, the presence of gender-specific differences in the presentation of DDs, and FASD as a commonly occurring DD. Ultimately, better support for children with DDs would reduce the burden of unmet needs for the children and their families, and for wider services.

Key words: SEND, ASD, ADHD, FASD, developmental disorders, EHCP
Introduction

Developmental disorder (DD) is a term used to cover a wide range of conditions all characterised by impairment or delay in the central nervous system (WHO, 2013). Three of the most common DDs are autism spectrum disorder (ASD), attention-deficit/hyperactivity disorder (ADHD) and fetal alcohol spectrum disorder (FASD). ASD conditions are characterised by deficits in social communication skills, unusual fixations and repetitive behaviours, and can be associated with intellectual disability (APA, 2013). ADHD conditions are characterised by reduced attention span, increased impulsivity and often hyperactive behaviour (APA, 2013). FASD is the name given to a range of conditions which can be caused by prenatal alcohol exposure (BMA, 2016). The neurological deficits seen in FASD can impact on attentional and social communication pathways, and, in severe cases, can lead to both ASD and ADHD as outcomes (Mukherjee et al., 2015).

Of the three conditions ADHD is thought to be the most common, with an estimated global prevalence of 2–7% (Sayal et al., 2018), and affecting between 1.4 and 8.8% of children in the UK (Polanczyk et al., 2007; Russell et al., 2014). ASD has an estimated global prevalence of 0.7% (Baxter et al., 2015) and is thought to affect 1.7% of children in the UK (Baxter et al., 2015; Brugha et al., 2016). Historically more males than females have been diagnosed with ASD and ADHD, at a rate of around 2.5:1 male to female (Whitlock et al., 2020; Zwaigenbaum et al., 2012), and there has been much debate about whether this is a true distribution or due to a lack of understanding of how these conditions present in girls and women (Allely, 2019). FASD is thought to affect 0.77% of children globally and 2% in Europe. Estimates for the UK are at 3.24% (Lange et al., 2017), while the first study to assess for FASD directly, in a small sample, suggests a conservative estimate of 1.8% (McCarthy et al., 2021). However, globally, only a small fraction of cases of FASD are diagnosed, and FASDs have been described as an under-recognised area of special educational needs (Blackburn et al., 2012; Carpenter et al., 2013). FASD training is needed across different professional groups, including teachers and other education professionals (Blackburn et al., 2012). The relatively low levels of training in the UK are reflected in significant knowledge gaps, and low levels of awareness and confidence among professionals (Mukherjee et al., 2015; Schölin et al., 2021).

The Assessing the Prevalence of Developmental Disorders in Greater Manchester Children Study (ADD-GM) (McCarthy et al., 2021) was an active case ascertainment study that aimed to estimate the prevalence of FASD and other DDs in mainstream schools in England. ADD-GM was
the first study of its kind in the UK and was part of the Greater Manchester Health and Social Care Partnership’s wider programme to prevent alcohol-exposed pregnancies (Reynolds et al., 2021). In ADD-GM, all children in year group 4 (aged eight or nine years) had been first pre-screened and had been invited for full assessment if they were: on the school’s special needs register; small for their age (and/or with a small head circumference); in the care of the local authority (or previously in care); or if parents or teachers were concerned about the child’s development or behaviour. Screen-positive children had been invited for neurodevelopmental assessments, including two hours of tests with children and a two-hour medical history interview with mothers. Parents were sent a comprehensive report indicating likely diagnoses of a neurodevelopmental condition, and suggestions for actions they could take to support their child, including onward referral. Assessments for ADD-GM had been completed just prior to the UK’s first national Covid-19-related lockdown (March 2020), which involved widespread disruption, including closing schools to most children and a move to online learning. This formed a significant backdrop to parents’ receiving the individualised report for their child. This study described here was carried out alongside ADD-GM, with the aim of exploring the experience of the parents of children who had a disorder identified by the screening study. As well as exploring feelings and opinions on the process and outcome, the study aimed to find out whether this had been impacted by Covid-19-related lockdowns.

Methods

Details of participants

Parents of children in two participant schools who had recently taken part in the wider ADD-GM study were invited to take part. After excluding one parent who had not yet received their child’s report, 16 invitations were sent. Contact was made via email or SMS text message (the parents’ preferred method of communication) to ask if they would be willing to give feedback on their experience of taking part in the ADD-GM study. Nine responded and received a participant information sheet and consent form, of whom six returned consent forms and agreed to take part in the interviews.

All participants (n = 6) were female, identified as White British ethnicity, spoke English at home, were born in the UK and were aged between 25 and 43 years (mean 34 years). Three of the participants were the child’s birth mother, one was an adoptive mother, one was a legal guardian and one was a
foster parent. Four parents had achieved higher-level education. The remaining two had achieved fewer than four GCSEs (or equivalent). The GCSE (General Certificate of Secondary Education) is an academic qualification obtained in England at the age of 16 years. Since most people sit nine or more GCSE examinations, achieving four or fewer is considered a low attainment.

Two of the parents had opted their child into the study because of concerns over the child’s development (‘parent opt-in’), one child had been invited due to being small for their age, two children had been invited due to being under the care of the local authority, and one who had previously been under the care of the local authority. Three of the parents had received reports stating that their child may have FASD, two received reports suggesting ASD and one received a report suggesting ADHD (table 1). All parents had received their child’s final report on the assessment at least 30 days previously to ensure parents had adequate time to read and digest the results.

Further information about participants and their children is given in table 1. Children are referred to using pseudonyms.

**Design**

A qualitative method was used in line with other studies examining parents’ experiences of having a child with a DD (Mohamed et al., 2020; Myers et al., 2009; Salmon, 2008). The interview guide comprised questions that probed the experience of taking part in the study. Parents were asked to describe their feelings, thoughts and experiences of having a child identified as having a DD during the course of the ADD-GM study. Interviewees were asked if their child had been seen by a specialist, and about the support their child had received from school before and after taking part in

<table>
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<tr>
<th>Pseudonym of child</th>
<th>Gender</th>
<th>Reason for invitation</th>
<th>Identified disorder</th>
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<tbody>
<tr>
<td>Alexander</td>
<td>Male</td>
<td>Prev LA</td>
<td>FASD</td>
</tr>
<tr>
<td>Charlotte</td>
<td>Female</td>
<td>Parent opt-in</td>
<td>FASD</td>
</tr>
<tr>
<td>Emily</td>
<td>Female</td>
<td>Parent opt-in</td>
<td>ASD</td>
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<tr>
<td>George</td>
<td>Male</td>
<td>LA</td>
<td>ADHD</td>
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<tr>
<td>Kyle</td>
<td>Male</td>
<td>LA</td>
<td>ASD</td>
</tr>
<tr>
<td>Lucy</td>
<td>Female</td>
<td>Small for age</td>
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the ADD-GM. To explore the experience of taking part, we asked questions such as: ‘How did you feel when your child was invited?’ and ‘What did you make of the results?’ The support received after getting the report was probed with questions such as: ‘Have you accessed any of the support services mentioned in your report?’, ‘Has your child been seen by a specialist?’ and ‘What impact has the report had on your child’s school?’ We also asked whether Covid-19 and the associated national lockdown had an impact (for example, ‘What impact has receiving this report had on your experience of lockdown?’ ‘How has lockdown impacted on your ability to access support?’ ‘What impact has lockdown had on your child’s behaviour?’). Questions were developed by the first author and the last author, based on existing literature and on the research questions.

Digital recordings were made of each interview, with a mean duration of 28:00 minutes (range: 14:00–37:00) and the interviews were transcribed verbatim by REM.

**Analysis**
Data analysis was informed by processes of thematic analysis – an inductive approach designed to identify, represent and report thematic patterns that occur within the data (Braun & Clarke, 2006).

As the research questions guided the study, the first level of qualitative data analysis for interview transcripts and observation data was *a priori*. It allowed analysis to organise, condense and categorise data. This was followed at the second level by an inductive process that allowed initial codes or sub-categories that described participants’ experiences to emerge. After initial codes had been identified in the first transcripts, subsequent codes could be compared and contrasted to identify similarities and differences in categories. Categories stayed close to the original expressions or records. Some were changed through abstraction and through combining of sub-categories during the analytical process (Charmaz, 2000). To increase validity, the transcriptions and thematic analysis that had been conducted by the lead author were checked and coded independently by a second researcher (the sixth author). The findings were then compared and reviewed by a third researcher (the second author).

**Ethics**
Informed consent, the right to anonymity and the right to withdraw at all stages of the research were established and re-confirmed prior to starting the
recording. Ethical approval was granted by the University of Salford, UK for the ADD-GM project (HSR1819-100 May 2019) and for this sub-study (HSR1819-100, May 2020).

Overview of findings
Analysis of the data revealed three main themes. One of the themes related to participation in a research study, and is not reported here as it is less relevant to education. Table 2 lists the themes and sub-themes reported in this article. The first theme was ‘barriers to support’. This was a strong theme since none of the children had previously received a neurodevelopmental diagnosis and therefore their needs had not been met at school. The second theme was ‘lockdown’ and was related to the unprecedented circumstances of the first UK national lockdown as a result of the Covid-19 pandemic. It was prompted by our specific questions about lockdown.

Findings: barriers to support
Universally, parents described obstacles to obtaining appropriate diagnosis, intervention, understanding or compassion for their child’s additional needs. This included failure to recognise and acknowledge the presence of additional needs, especially in school. Parents described resistance from the school when communicating their concerns about their child’s development and/or actively advocating for formal assessment or support. This appeared to be a particularly significant issue where the children were described as quiet, well-behaved girls. There was overlap here with the theme of lockdown, whereby parents described current barriers to diagnosis and support rather than those they had experienced in the past.

Table 2: Themes identified using framework analysis

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<th>Theme</th>
<th>Sub-theme</th>
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<td>Barriers to support</td>
<td>Accessing support from school</td>
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<td>Well-behaved girls</td>
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<td>Accessing support during lockdown</td>
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<td>Shame and stigma as a barrier to participation</td>
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<td>Lockdown</td>
<td>Child’s experience</td>
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<td>Accessing support during lockdown</td>
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<td></td>
<td>Shame and stigma as a barrier to participation</td>
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<td>Participation in a research study (not included in this article)</td>
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‘Brick walls’: accessing support from school

Several parents had experienced resistance when communicating with schools and felt their concerns about their child’s development had been dismissed. In the wider ADD-GM study, being on the school’s register for special educational needs was one of the screening criteria for full assessment. However, none of the six parents who came forward for this study had their child listed by the school as having special educational needs. The reasons that their children had been assessed was because they were opted in because of parental concerns, had been invited due to being currently or previously in the care of the local authority, or because they were small for their age. When talking critically of the response from their child’s school, parents spoke in a manner which suggested they were choosing their words carefully and were cautious with the tone they used. Despite this, parents described significant resistance from schools when they raised their concerns:

‘I spoke to the teachers at parents evening first and I’d expressed our concerns to them … and both the teachers were there, and they said, “Err yeah but well we don’t see that little girl”, and that had been the same response I’d had previously from the teacher last year.’ (Emily’s mother)

Alexander’s mother reported that home-schooling during lockdown had made it possible to see more of Alexander’s and other children’s work. This parent felt there was an obvious difference, enough to suggest her child was significantly behind his peers and this was at odds with communication from the school, which had assessed the child as achieving age-appropriate levels:

‘So yeah, as a result of home schooling I think I’ve got a better idea of where Alexander is because they [school] say he’s doing OK but, well, school are telling me he’s where he should be and … I’m not criticising them … but I think he’s quite behind. So, it’s quite an eye opener and you shouldn’t compare, but I see the other work that the other kids are doing, assuming the parents aren’t doing it, he’s quite far behind that.’

Lucy’s mother reported struggling to access support from school for her child for two years. She described how the school argued against her request for assessment for her daughter, denying her concerns, and how she was left feeling patronised and belittled. This parent also used the
metaphor of a brick wall for trying to get the school to acknowledge that Lucy was struggling:

‘yeah, about two years I’d asked for help. I’d asked to have her assessed and they’d been like there’s no problem there’s no problem and … clearly there is. Yeah. I’m always up against a brick wall with them because they just look at me like I’m stupid, I’m not stupid…’

Lucy’s mother went on to explain that she felt the school was reluctant to acknowledge an issue and pursue assessment if they felt they were unlikely to get extra financial support for the child in the form of an Education, Health and Care Plan (EHCP):

‘I’m not gonna lie I’ll even say it to the school’s face, unless they know that child’s gonna get the extra funding they don’t put anything through.’

For the same child, while school staff were reluctant to refer, they were also reluctant to allow adaptions without a diagnosis, and Lucy’s mother described with frustration the process of trying to get support for her daughter as circular:

‘I’ve mentioned can she take ear defenders in, but they say no ‘cos she’s not diagnosed with anything … it’s a circle’.

Another parent found that school staff now having awareness of her child’s additional needs has been hugely beneficial to her daughter’s experience at school. This is despite the changes only being slight, such as letting the child eat lunch in a quiet room and encouraging the use of sensory toys in the classroom.

‘they’ve put lots of little, just sort of little tweaks in place … the teacher will kind of say “change your pen” and direct her to the pen that’s got the chewy buddy on’ (Emily’s mother)

‘Little girls are getting missed’: quiet well-behaved girls
A strong sentiment expressed in interviews with the parents of girls was the idea that their child’s extra needs had gone undetected or unacknowledged by the school as they were quiet and well-behaved. Gender was not mentioned
by the parents of boys. By comparison, the parents of girls described having more contact with the school. They were more likely to have initiated contact with the school themselves and expressed more frustration with the process than the parents of boys:

‘I think a lot of little girls are getting missed because they do try, and they are more aware of what’s socially expected of them even if they don’t always enjoy it.’

(Emily’s mother)

‘They say “yeah [mother’s name] she is quiet in class; she is a bit behind her peers”. Well DING DONG! Why is that bell not being rang? … but it tends to be the child who is being very disruptive in class, they will go for that child first rather than thinking Lucy is very quiet and wondering why’.

(Lucy’s mother)

Emily’s mother felt having the support of other girls in Emily’s circle of friends was masking some of Emily’s symptoms. This parent implied that this would not have happened with a circle of friends who were boys and that this could have been one explanation as to why the school had not acknowledged her daughter’s additional needs earlier:

‘one of the things that has been picked up on while we’re having the meetings is that sometimes they [peers] can step in and kind of tell Emily what she needs to do to. Y’know that rather than letting her experience it, they say “Emily you don’t like this, you need to do this, or you need to do that”.

‘Not enough evidence’: accessing further support
Parents were asked what had happened since receiving the diagnostic findings and suggestions from their child’s participation in the wider ADD-GM study to identify DD, and their responses highlighted further barriers to accessing support. This included struggling to arrange referral and diagnosis and their feelings about accessing parent support groups and advocacy services.

Barriers ranged from resistance from schools and medical professionals to local variations in referral pathways and their own readiness to ask
for a referral. Some parents had been referred but none had received a confirmation letter or appointment date. One parent described being sent back and forth between the school and the family doctor. The way Lucy’s mother recounted the scenario communicated feelings of frustration, disbelief and injustice:

‘I’ve been to doctor, they said school need to refer to CAMHS and basically went back to school, told school doctors need school to refer, they say we don’t have enough evidence, I said well she been in your school since nursery. You must have something on her and basically and went back to the doctors and said “Look, one of yous is going to refer her!”’

Charlotte’s mother described how her family doctor had asked for a written list of symptoms as well as the study report findings before making any referral for a child who had been identified by the ADD-GM study as having FASD:

‘She [the family doctor] said to write some notes of my experiences, y’know symptoms that I’d noticed at home and send in a copy of your report with my notes. But I haven’t written up my notes yet, but she thought without that, like some notes from me as well that she wouldn’t … was unlikely she’d get a referral for … anywhere really.’

Most parents talked about referral as an option or possibility rather than the immediate or obvious next step. They talked with reticence and often sounded unsure and vague when discussing it. George’s mother, who is a foster carer, reported that:

‘it was brought up at the LAC [looked-after child] review and they’re not sure what they’re gonna do about it at the moment whether we pursue it with him being assessed for this condition that he might have. So, we just talking about that at the moment.’

Kyle’s mother described trying to access mental health services for her child and expressed her annoyance at reaching out for help for her child, only to be offered parent courses:
'the community mental health team said they don’t work with children under the age of eleven … so, face to face anyway it’s more erm doing courses for the parents or carers … and we’ve done all the courses!’

All parents were sent information leaflets with their child’s results, including details of local support groups, but none reported having contacted the support organisations listed on them. Lucy’s mother described why she felt a support group was not helpful for her, as most of her child’s issues were school-related. The same parent did indeed report improvement in her daughter’s symptoms as a result of being home-schooled during lockdown:

‘there is some good support groups local up here that you can connect to but … because going to support groups doesn’t help in school. when I’m trying to help Lucy in one way, she’s in school nine to three [9am–3pm], it’s all irrelevant, because school are gonna do what they’re gonna do’.

Findings: lockdown
Government-mandated lockdown restrictions related to Covid-19 between March and June 2020 were in force during the time in which the interviews were conducted, and this was a prominent theme in parents’ responses. Although schools were not open to all students during this time, some of the children in this study were still attending school either because, as per the UK Government rules at the time, they were under the care of the local authority, or because their parents were ‘critical workers’ (that is, those working in health and social care, and key public services).

‘Losing routines and bringing families closer together’: the child’s experience
Overall, participants reported that lockdown had been a positive experience for their child and few reported increased anxiety or sleep or eating disturbances.

For children identified by the study as having ASD, parents reported that their children liked the increased time at home and reduced travel and social expectations, such as for food shopping or visiting relatives. Parents expressed some surprise and relief at this:
‘She’s quite enjoyed lockdown, because Emily doesn’t like going to crowded places, she doesn’t like going [to the] supermarket, going shopping … and she doesn’t particularly like going visiting family members.’

(Emily’s mother)

Parents of children identified as having FASD all reported a change in behaviour under lockdown. Although it had not been immediately obvious, as time went on their child had struggled more with the loss of routine and social interaction. This included issues with behaviour and emotion regulation, impulse control, and feelings of shame. One parent referred to their child’s lying (which can be a symptom of poor memory function in those with FASD, which is more accurately described as ‘confabulation’). Some of these changes in behaviour are exemplified by the following comments:

‘We’re struggling with him; his attention span isn’t great, he gets very frustrated, he gets angry constantly.’

(George’s mother)

‘She handled it really well at first. For the first four weeks, y’know she was handling it really well with doing her schoolwork but once the novelty has worn off at home her behaviour’s kinda come back again.’

(Charlotte’s mother)

‘I think from a behaviour perspective yeah we’ve had a lot of tears, a lot of tantrums and he’s quite down on himself … we’re getting a lot of “I’m a terrible child” and I tell you what else has increased, the lying.’

(Alexander’s mother)

Several of the children had continued to attend school through lockdown and parents spoke about this positively. They felt this was helpful to their child’s well-being. Some parents reported that still being able to attend school kept in place the structure that school provides day to day in terms of dictating a fixed time to get up and a regular mealtime, but also by preserving the structure of a working five-day week with a two-day weekend. Kyle’s mother commented, ‘he’s still been attending school so; he has been kept in the same routine which I think helps’.

No parents reported increased anxiety or fears in their children specifically related to Covid-19. Emily’s mother said:
‘With regards to the news she doesn’t know that much, she knows that it’s a virus and it’s out there, there and that’s the reason mummy and daddy don’t go to the supermarket together anymore and the reason we don’t see family members.’

Several parents reported extra time together as a family unit as a positive aspect of lockdown and talked about the benefits of extra time at home. Some of these benefits included learning life skills that they would not have had the opportunity to learn about at school, for example, about baby care:

‘It’s really nice. They’ve been with us more … er … I think as a unit … and they’ve been helping out with the baby a lot, read to him and help with bathing and things like that so they’re learning a lot. Yeah, in that respect I think it’s brought us closer together’

(Kyle’s mother)

Only one parent reported concerns that their child’s education had been set back by lockdown; others reported that their child was missing friends and extracurricular activities:

‘She’s gonna be a year behind as she missed so much school and I know she’s gonna struggle because she’s definitely going to need extra help when she goes back.’

(Lucy’s mother)

‘he’s saying he’s missing his activities, his football cricket and swimming, but we all are missing the things we can’t do. We’ll probably all appreciate it more when we go back and do those things.’

(Kyle’s mother)

‘Waiting ’til we come out of lockdown’: finding support during lockdown

Many participants referred to services not being available or being further delayed during lockdown, and there was a sense of stoicism and acceptance that nothing would be done until lockdown finished. This was in contrast to the frustration that was felt by those who had perceived that schools had failed to detect a potential problem:

‘so we’ve had a referral to the mental health team for her so at the moment it doesn’t seem that … but they’re not working with families at the moment and its more parent and carers doing courses, well we’ve been on
courses! So, there’s not really a lot at the moment that can be done ’til we come out of lockdown’

(Kyle’s mother)

In the case of Lucy’s mother, the end of lockdown and return of ‘normality’ was going to be the point at which she would vigorously pursue avenues of support for her daughter:

‘When this corona goes, if it goes and life goes back to some kind of normality. I’m not even gonna lie, I’m gonna be on the phone, I’m gonna be badgering them all.’

(Lucy’s mother)

The waiting time for referrals had also delayed school-level support, as one parent felt they needed to wait for a referral before sharing their child’s report with the school. Other parents mentioned that they had not accessed support groups due to lockdown. However, in contrast to the previous distress and frustration caused by resistance from school or general practitioners prior to lockdown, the lack of progress due to lockdown seemed to be accepted. There was a sense of being in stasis at this unique moment in time, as the following comments show:

‘I was waiting to get referred erm and get a bit more information before I sort of jumped in and started talking to school.’

(Alexander’s mother)

‘Obviously I haven’t accessed that [support groups] as all the children are off school and I’m home schooling.’

(George’s mother)

Discussion
This study reports on parents’ experiences of school support for children with a previously undiagnosed developmental disorder (DD) whose difficulties had previously been unrecognised by the school. Although parents of children on the special educational needs and disability (SEND) register had also been invited to take part, the children of those parents who volunteered to be interviewed were not on the schools’ register. All the children whose parents took part had experienced difficulties in engaging with learning at a level that could reasonably be expected to have been detected by a mainstream school in children of this age. Not surprisingly, therefore, a major theme was frustration that the school had not detected and supported their children.
Failure to detect and support

Parents reported many barriers to accessing the correct support for their child, expressing frustration and distress. Schools failing to detect that a child may need assessment or extra support increases the risk that the individual will present to services later with complex secondary mental and physical health issues that may have been preventable (APA, 2013; Streissguth et al., 2004). Long-term lack of support can lead to increased risk of interaction with the criminal justice service, children’s services and hospital admissions (Bradshaw et al., 2015; Streissguth et al., 2004). Therefore, failing to detect and support DDs is likely to have significant multi-service cost implications in the long term. In two cases, parents described having had their concerns dismissed by the school for some time. The child we have called Lucy was found to have a Full Scale IQ score of 66 (Canivez et al., 2017). While not a complete picture of a child’s abilities, it does indicate a strong possibility that a learning disability is present. This had not been detected by the school despite Lucy’s parent asking for assessment and additional support for two years. The issue was raised particularly for children who were described as ‘quiet, well-behaved girls’, and this emerged as a sub-theme in our analysis. As observed by Lucy’s mother, children who may be perceived by their schools as disruptive are more likely to be assessed than well-behaved children, and this is reflected in the findings of previous research (van Bergen et al., 2015).

Similarly, Emily’s mother described raising concerns over a year prior to taking part in the research. Emily’s mother attributed the failure to detect additional needs in girls to masking of symptoms. This is consistent with the findings of previous studies which have explored ASD in females (Allely, 2019; Hull et al., 2020). An increasing number of studies have found that ASD is underdiagnosed or unrecognised in females (Russell et al., 2011; Whitlock et al., 2020). One of the potential explanations for this is camouflaging, whereby females with ASD compensate for and mask their autistic characteristics (Dean et al., 2017; Hull et al., 2020; Halsall et al., 2021; for review, see Tubío-Fungueiriño et al., 2021). Camouflaging strategies are diverse. Some can be very simple (for example, when someone develops rules for using eye contact in a conventional manner) and others can be more complex. An example of a complex camouflaging strategy would be a teenage girl with autism who purposefully studies the behaviour of a girl without autism at school, and then, over time, adopts her attitudes, dress, gestures and facial expressions – essentially imitating the behaviour of the girl without autism. By doing this, the individual develops a persona in
order to enable them to navigate social situations (Mandy, 2019). Given
the well-established finding of camouflaging in many females with ASD,
teachers and other professionals should receive training on and have an
awareness and understanding of how some girls with ASD may engaging
in camouflaging behaviours.

In the UK, the updated Special Educational Needs Code of Practice (DfE,
2015; Ponsford et al., 2017) introduced the new Education, Health and
Care Plan (EHCP) for children with SEND. The intention of these reforms
was to ensure integration between educational provision and training pro-
vision, and health and social care provision, where this would promote
well-being and improve the quality of provision for disabled children and
young people and those with SEND. The most recent data for England
show that although 12.8% of primary school children have acknowledged
special educational needs, fewer than a third have been given EHCPs, sug-
gest issues in accessing the new EHCPs except for children with the
most complex SEND. Boys are more likely to be classed as having SEND
(64.6%) and even more likely to be given an EHCP (ONS, 2020; Ward &

The account given by Charlotte’s mother, who took their report to the
family doctor only to be asked to make notes about Charlotte’s symptoms
before a referral could be made, illustrates a significant barrier to accessing
support, which has the potential to increase health inequalities, and echoes
the findings of other research (Petrenko et al., 2014). This child was sus-
pected of having FASD, and this barrier might have arisen because there
is currently no official diagnostic pathway for FASD (although this is in
development in England: NICE, 2021). This highlights the importance of
increasing awareness of FASD and suggests there is a need to improve the
training offered to professionals. Relative to other conditions, FASD is
associated with higher levels of shame and stigma (Mukherjee et al., 2015;
Salmon, 2008). This could mean parents are less likely to pursue diagnosis
and support. Indeed, Charlotte’s mother came across as feeling defeated
by the system and was yet to write up notes on her daughter’s symptoms
for their family doctor. By comparison, in this study, parents of children
identified with ASD and ADHD were more comfortable naming and dis-
cussing these conditions, which are more widely known and understood
in recent years. This suggests that drawing on the success of campaigns to
increase awareness of ASD and ADHD could bring about similar progress
for FASD and other DDs.
The study was carried out at a particular moment in time when the UK had adopted the first national Government-mandated Covid-19 restrictions. There was an acceptance that there was going to be a longer wait for services, and a reluctance to push for a referral. This may have been because participants were aware of the general pressure on the health care system, and that waiting lists were longer for care for many different conditions. The feeling that everyone is having to wait seemed to be linked to a degree of stoicism and acceptance.

**Lockdown**

The wider study to identify DDs in schools, the ADD-GM study, was in the process of completion by the time the first UK national lockdown (between March and June 2020) was implemented. One research aim was to investigate how this unprecedented situation had interacted with the experience of receiving the report that stated that their child had a previously unrecognised DD. While schools were closed for the majority of pupils, some of the children in this study were still attending school, either because they were deemed ‘vulnerable’ (due to being in the care of the local authority) or because their parents were ‘critical workers’ (for example, health care or key public service workers).

Overall, the parents reported little negative impact of lockdown on their children, whether children were still attending school or not. Parents of the children identified as having FASD were more likely to describe lockdown-related changes in behaviour, which could indicate increased levels of stress or distress in these children. Parents of children identified as having ASD felt their children benefited from reduced social interactions and travel. There is little research to date relating to children’s experiences of the first lockdown. Further research is needed to identify whether these positive stories are generalisable for parents of children with ASD or DDs or the general population, and whether the negative stories are generalisable to children with FASD.

Data collected by the Children’s Commissioner in England suggest that, in general, children’s levels of anxiety have increased as a result of Covid-19 (Children’s Commissioner, 2020). However, no parents in our study reported increased anxiety in their children related to awareness of Covid-19 or to media coverage of the pandemic. This is an interesting finding as anxiety is a symptom associated with all three conditions. It also contrasts with symptoms of anxiety that parents described in their children relating to attending school, travel and social interactions.
Strengths and limitations
Although over a third of the eligible parents took part in the study (six out of 16), the sample size was nevertheless small, making it difficult to extract findings related to factors such as the specific DD, children’s gender or parents’ socio-economic status. The sample size is adequate according to recommendations for the thematic method of analysis (Braun & Clarke, 2013). The findings of this study may be affected by the bias of the individual parent, or the interviewer. Transcriptions and analysis were repeated independently by another researcher and reviewed by a third in an attempt to improve the validity of the findings. The fact that parents of children already on the SEND register chose not to take part suggests that this sample may have been more reflective of those who were grateful to receive our report on their child’s DD. As with much qualitative research of this nature, it is possible that individuals with strong feelings were more likely to take part. These are limitations inherent to qualitative research and mean the results may not be generalisable to the wider population. Nevertheless, studies to investigate the prevalence of DDs in school settings are rare, and this study capitalised on an opportunity to explore parents’ experiences of taking part, thus making a novel contribution to the literature.

Conclusion
A wider study into the prevalence of DD, the ADD-GM study, provided a unique opportunity to examine the experiences of parents of previously undiagnosed children. The findings of this research revealed significant barriers for parents when accessing support for children with the three most common DDs. The mainstream schools that took part in the ADD-GM had failed to detect and support significant additional needs in this particular group of children. A lack of recognition was perhaps unsurprising for FASD, which, although common, is widely unrecognised. However, similar issues were raised by parents of children with ASD. Parents’ reluctance to discuss FASD may indicate that the shame and stigma associated with FASD is another barrier to detection, and it is likely to be another barrier to accessing support. Another barrier was experienced for parents of girls: for children described as ‘quiet, well-behaved girls’, parents reported that their concerns had been dismissed by the school for some time prior to their participation in the study. Parents felt that children with externalising behaviours were more likely to be assessed and supported, and that camouflaging of ASD symptoms in girls may be another factor affecting detection in mainstream schools.
The disruption caused by the Covid-19 pandemic was another barrier to accessing services. The national lockdown was a strong theme in itself, with experiences being markedly different between participants, from those whose children were still attending school, to those who were not, and for those whose behaviour improved during lockdown (typically for ASD) to those whose behaviour became worse (typically for FASD).

Finally, further research is needed to measure the extent of undetected and unsupported additional needs in primary schools in England.

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