Chromosomal microarray analysis in a cohort of underrepresented population identifies SERINC2 as a novel candidate gene for autism spectrum disorder

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Chromosomal microarray (CMA) is now recognized as the first-tier genetic test for detection of copy number variations (CNVs) in patients with autism spectrum disorder (ASD). The aims of this study were to identify known and novel ASD associated-CNVs and to evaluate the diagnostic yield of CMA in Thai patients with ASD. The Infinium CytoSNP-850K BeadChip was used to detect CNVs in 114 Thai patients comprised of 68 retrospective ASD patients (group 1) with the use of CMA as a second line test and 46 prospective ASD and developmental delay patients (group 2) with the use of CMA as the first-tier test. We identified 7 (6.1%) pathogenic CNVs and 22 (19.3%) variants of uncertain clinical significance (VOUS). A total of 29 patients with pathogenic CNVs and VOUS were found in 22% (15/68) and 30.4% (14/46) of the patients in groups 1 and 2, respectively. The difference in detected CNV frequencies between the 2 groups was not statistically significant (Chi square = 1.02, df = 1, P = 0.31). In addition, we propose one novel ASD candidate gene, SERINC2, which warrants further investigation. Our findings provide supportive evidence that CMA studies using population-specific reference databases in underrepresented populations are useful for identification of novel candidate genes.

Autism spectrum disorder (ASD) is a complex neurodevelopmental disorder characterized by impairments in social interaction and communication, as well as stereotyped behaviors and restricted interests. For approximately 80% of cases, the cause of ASD is still unknown. Due to clinical diagnostic criteria of ASD with broader spectrum of symptoms, the prevalence of ASD is increasing worldwide. The estimated ASD prevalence in the US was 1 in 150 children in 2000, but by 2012 that had increased to 1 in 68 children1. In Thailand, the incidence rate...
of severe ASD was estimated at 9.9 per 10,000 in children aged 1–5 years. Up to 70% of individuals with ASD also have intellectual disability/developmental delay (ID/DD). Genetic causes, including chromosomal abnormalities, copy number variations (CNVs), and single nucleotide variants are recognized as causative in 10–20% of ASD cases. Knowing the genetic cause of ASD can help with genetic counseling and evaluation of recurrence risk in the families.

Since 2010, chromosomal microarray (CMA) has been recommended as the first-tier clinical diagnostic test for detection of CNVs in patients with ASD, ID, DD and multiple congenital abnormalities (MCA) of unknown causes. CMA has a much higher diagnostic yield (10–20%) for these individuals than conventional cytogenetics (~3%). Most of CMA studies in these disorders have largely focused on Caucasian populations. To date only a few non-Caucasian studies have been reported. In our earlier study, we have categorized reference CNVs of more than 3,000 Thai individuals and created a CNV database that would facilitate the accurate clinical CNV interpretation for Thai patients. In this study, we aimed to screen and identify known and novel CNVs associated with ASD in a large cohort of Thai patients with ASD. This is the first report demonstrating the utility of CMA for detection of CNVs in Thai patients referred with ASD of unknown cause. Our findings also provide supportive evidence that CMA studies in underrepresented populations can be useful for identification of novel candidate genes.

## Results

### Diagnostic yield of CMA in Thai patients with ASD.

In our cohort of 114 Thai patients with ASD, a total of 742 CNVs was identified from all patients, ranging between 1 to 22 CNVs per patient. The size of the CNVs ranged from 1,632 bp to 18.9 Mb, with a median size of 86 kb. Among these CNVs, pathogenic CNVs and VOUS were identified in 7 (6.1%) and 22 (19.3%) patients, respectively. The overall diagnostic yield of CNVs in our cohort was 25.4% (29/114). The 29 patients with pathogenic CNVs and VOUS included 15 of 68 (22%) patients in group 1 and 14 of 46 (30.4%) patients in group 2. The summary and schematic diagrams of CMA results in the study are shown in Table 1, Figs 1 and 2. Although a higher frequency of both pathogenic CNVs and VOUS were identified in group 2 patients (30.4%) with the use of CMA as the first-tier test compared with patients in group 1 (22%) with the use of CMA as the second line test, no statistically significant differences were found in the frequencies between the 2 groups (Chi square = 1.02, df = 1, P = 0.31).

Parental testings were available for investigation of the inheritance status in 23 of 29 patients. Of these 23 patients, 5 de novo CNVs, 11 maternally inherited CNVs and 7 paternally inherited CNVs were identified (Fig. 2b). However, parental samples were not available in 3 pathogenic CNVs and 3 VOUS which was a limitation.

|                      | Group 1 (Retrospective cases) | Group 2 (Prospective cases) | Total | Chi squared test (P-value) |
|----------------------|-------------------------------|----------------------------|-------|---------------------------|
|                      | N = 68                        | N = 46                     | N = 114|                           |
| Pathogenic CNVs and VOUS | 15/68 (22%)                  | 14/46 (30.4%)              | 29/114 (25.4%) | 0.31                     |
| Pathogenic CNVs and VOUS, likely pathogenic CNVs | 9/68 (13.2%)                | 5/46 (10.9%)              | 14/114 (12.3%) | 0.71                     |
| Pathogenic CNVs | 5/68 (4.4%)                | 4/46 (8.7%)              | 7/114 (6.1%) | 0.35                     |
| Deletion | 1                           | 3                        | 4     | —                        |
| Duplication | 2                           | 1                        | 3     | —                        |
| VOUS | 12/68 (17.6%)                | 10/46 (21.7%)             | 22/114 (19.3%) | 0.59                     |
| VOUS, likely pathogenic CNVs | 6/68 (8.8%)                | 1/46 (2.2%)              | 7/114 (6.1%) | 0.15                     |
| Deletion | 2                           | 1                        | 3     | —                        |
| Duplication | 4                           | 0                        | 4     | —                        |
| VOUS, likely benign CNVs | 3/68 (4.4%)                | 2/46 (4.3%)              | 5/114 (4.4%) | 0.99                     |
| Deletion | 2                           | 1                        | 3     | —                        |
| Duplication | 4                           | 0                        | 4     | —                        |
| VOUS (no subclassification) | 3/68 (4.4%)                | 7/46 (15.2%)             | 10/114 (8.7%) | 0.04                     |
| Deletion | 1                           | 2                        | 3     | —                        |
| Duplication | 2                           | 5                        | 7     | —                        |
| Benign | 53/68 (78%)                 | 32/46 (69.6%)             | 85/114 (74.6%) | 0.31                     |

Table 1. Comparison of clinical CNVs identified in the two groups of our ASD subjects. The group 1 patients were first evaluated for the common genetic causes of ASD and all patients had normal karyotype and negative DNA tests for Fragile X syndrome and MECP2 mutations. Therefore, group 1 represented the ASD patients with the use of CMA as a second line test and group 2 represented the ASD patients with the use of CMA as the first-tier test. VOUS, Variant of uncertain clinical significance; *P-values were calculated using Chi-square test using GraphPad QuickCalcs (https://graphpad.com/quickcalcs/contingency1.cfm).
of our study. The clinical characteristics of patients and details of the pathogenic CNVs and VOUS are summarized in Table 2.

**Rare inherited CNVs and known syndromes.** 18 of the 23 CNVs in this study were transmitted from an asymptomatic parent. Most of the VOUS with inherited CNVs were below 1 Mb, but five CNVs were more than 1 Mb in size including four duplications and one deletion. Interestingly, one (TM54-3) was a 1.49 Mb deletion encompassing 2 OMIM genes (*PTP4A3*, *GPR20*), which was inherited from an asymptomatic father. Deletions in these regions have been reported as both pathogenic CNVs and VOUS in the ISCA/ClinGen and DECIPHER databases. Thus, it is difficult to interpret the significance of the deletion in this patient. However, we assumed that it was a VOUS, because genes in this region were not involved in neurological function which may have been associated with the phenotype of this patient.

Within the group of patients with pathogenic CNVs, there were patients with well-known microduplication or microdeletion syndromes, including the 1q21.1 duplication, 15q13.3 microdeletion, 16p13.11 microduplication, 22q13.3 deletion (Phelan-McDermid syndrome), 18q distal deletion, and 4p16.3 distal deletion (Wolf-Hirschhorn syndrome). Three of the pathogenic CNVs were inherited from a healthy parent. This is in line with the fact that some of these established syndromes are known to have incomplete penetrance and/or variable expressivity.

**De novo CNVs and novel candidate genes for ASD.** Among the *de novo* CNVs, one was a large deletion associated with the known 9q21.13 microdeletion syndrome, while the other 4 *de novo* VOUS were rare duplications of 127–220 Kb in size. A 127 kb duplication at 2q14.1 overlapping the *DPP10* gene was identified in a female proband (TM18-3) and a 159 kb duplication at 3p26.3 overlapping *CHL1* gene was identified in a male proband (TM4-3). Both the *DPP10* and *CHL1* genes have been previously reported as candidate genes of ASD. One male patient with non-syndromic ASD (TM41-3) had a 220 kb duplication at 1p35.2 including the *FABP3*, *SNRNP40*, *NKAIN1* and *SERINC2* genes. A *de novo* 217 kb duplication at 18q22.3 including the *ZNF407*, *CNDP2* and *CNDP1* genes was identified in a male proband with non-syndromic ASD (AR12-3). Table 3 provides details.
Incidental findings of common α-thalassemia mutations. In this study, we identified the deletions of α-globin genes, HBA1 and HBA2 as the most common incidental findings in our patient cohort. Chromosomal microarray was able to detect the common deletions of HBA1 and HBA2, leading to the identification of α-thalassemia deletions in 9 of the 114 ASD patients (7.89%). These α-thalassemia deletions were confirmed by single-tube multiplex-PCR assay as previously described35. Among the 9 patients, eight had available DNA for validation. The results confirmed 5 cases with heterozygous alpha-thalassemia-1 (–α3.7/αα), one case with heterozygous alpha-thalassemia-2 (-αα/αα), one case with homozygous alpha-thalassemia-2 (-αα/αα) and the other case with compound heterozygous alpha-thalassemia-1 and alpha-thalassemia-2 (-αSEA/αα), also known as Hb H disease (Supplementary Table S2).

Detection of absence of heterozygosity (AOH) regions. We also analyzed the AOH regions greater than 10 Mb in size in all patients. One case showed two large AOH regions of 40.6 Mb (15q11.1-15q22.2) and 12.4 Mb (15q26.1-15q26.3) interrupted by a region of heterozygosity (Supplementary Figure 2), a condition which was suspected to be from uniparental disomy (UPD). Methylation-specific polymerase chain reaction (MS-PCR) analysis of SNRPN gene was then performed using primers as previously described36,37, and the results confirmed that this patient had maternal uniparental isodisomy of the 15q11.2 region resulting in Prader-Willi syndrome (Supplementary Figure 3). Additionally, multiple large AOH regions (average 158.02 Mb) were also detected in another patient, encompassing 5.84% of the genome (data not shown), suggesting a consanguineous marriage between first cousin parents (coefficient of inbreeding (F) 1/16). Since this patient had an increased chance of having an autosomal recessive disease, it might be beneficial to further search for homozygous gene mutations by means of whole exome sequencing.

Discussion

We performed CMA in 114 Thai patients with ASD of unknown cause, resulting in an overall diagnostic yield of 25.4% composed of both pathogenic CNVs and VOUS. The diagnostic yields of only pathogenic CNVs and of VOUS alone are 6.1% and 19.3%, respectively. Our pathogenic CNV's diagnostic yield is in line with those of other CMA studies in non-Asian patients with ASD, which range from 5.4% to 8.6%9,12,18,22. However, it is difficult to directly compare such diagnostic yields among studies due to the use of different inclusion criteria for diagnostic yields. Some studies report diagnostic yields from only pathogenic CNVs18,11,13–18,20,22,24,26,28,29,32, while others include combined pathogenic CNVs and likely pathogenic CNVs10,23,25,26 or combined pathogenic CNVs and VOUS12,19,21,27. In addition, previous studies have had different inclusion criteria for participating patients.
| Case ID | Sex | Clinical diagnosis | Additional clinical features | Chromosome region | Coordinates (hg19) | Gain/loss | Size | Gene involved* | Inheritance |
|---------|-----|-------------------|-----------------------------|------------------|-------------------|-----------|------|----------------|-------------|
| A. Pathogenic CNVs (N = 7 of 114) |
| AR82-3 | M   | Non-syndromic ASD | Macrocephaly 1q21.1-1q21.2 | chr1:146571244-147825548 | Gain 1.25 Mb | 9 OMIM genes | Maternal |
| TU22  | M   | ASD, ID, seizures | Microcephaly 4p16.3 | chr4:49450-199778 | Loss 1.95 Mb | 22 OMIM genes | NA |
| 2715   | M   | ASD, ID, seizures | Dysmorphogenic features 9q21.11-9q21.2 | chr9:7206666-79948984 | Loss 7.88 Mb | 22 OMIM genes | De novo |
| AR83-3 | F   | Non-syndromic ASD | None 15q13.2-15q13.3 | chr15:30371774-32514341 | Loss 2.14 Mb | 8 OMIM genes | Maternal |
| TM50-3 | M   | Non-syndromic ASD | None 16p13.11 | chr16:15129970-16633361 | Gain 1.50 Mb | 11 OMIM genes | Paternal |
| TU17   | F   | ASD, ID, MCA | Microcephaly, cleft lip and palate, absent philtrum, abnormal ear canal, hypertelorism, flat nasal root, prominent nose 18q21.33-18q23 | chr18:59030666-78015180 | Loss 18.9 Mb | 41 OMIM genes | NA |
| 2950   | F   | ASD Familial history of lissencephaly | 22q13.33 | chr22:50537901-51211392 | Loss 673 kb | 23 OMIM genes | NA |

*B: VOUS, likely pathogenic CNVs (N = 5 of 114)  
| TM41-3 | M   | Non-syndromic ASD | None 1p35.2 | chr1:31707971-31928387 | Gain 220 kb | FABP3, SNRNP40, NKAIN1, SERINC2 | De novo |
| AR44-3 | M   | Non-syndromic ASD | Macrocephaly 1q42.2 | chr1:231763496-231877073 | Loss 113 kb | DISC1 | NA |
| TM18-3 | F   | Non-syndromic ASD | None 2q14.1 | chr2:1116.548,326-116,675,350 | Loss 127 kb | DPP10 | De novo |
| TM4-3  | F   | Non-syndromic ASD | None 3p26.3 | chr3:104972-264293 | Gain 159 kb | NA | NA |
| AR33-3 | M   | Non-syndromic ASD | Abnormal hearing test 17p13.3 | chr17:228978-440497 | Gain 211 kb | NA | NA |

*C: VOUS, likely benign CNVs (N = 5 of 114)  
| TM13-3 | M   | Non-syndromic ASD | Macrocephaly 4q24 | chr4:103560963-103658350 | Loss 97 kb | MANBA | Paternal |
| PS37-3 | M   | Non-syndromic ASD | seizures 10q23.1 | chr10:84017608-84115017 | Loss 97 kb | NRG3 (intronic) | Maternal |
| AR33-3 | M   | Non-syndromic ASD | None 17p13.3 | chr17:228978-440497 | Gain 211 kb | FAM101B, VPS35 | Paternal |
| RA29   | M   | ASD, ID | None 15q13.3 | chr15:32,389,362-32,415,088 | Loss 25 kb | CHRNA7 | NA (Not paternal) |
| AR12-3 | M   | Non-syndromic ASD | Abnormal hearing test 18q22.3 | chr18:72186404-72404396 | Gain 217 kb | ZNF407, CNDP2, CNDP1 | De novo |

*D: VOUS (no subclassification) (N = 10 of 114)  
| AR91-3 | M   | Non-syndromic ASD | None 2q13 | chr2:112581082-112858485 | Gain 277 kb | ANAPC1, MERTK | Maternal |
| RA15   | M   | ASD, ID | None 2q12.2-2q12.3 | chr2:106637427-107642482 | Gain 1.0 Mb | C2orf40, UXX1, RLPLA, BGP33, ST6GAL2 | Paternal |
| TM35-3 | F   | Non-syndromic ASD | None 4q28.2 | chr4:128901402-130163564 | Gain 1.26 Mb | SCL11, PHF17, PGRMC2 | Paternal |
| 2546   | M   | ASD, GDD, MCA | Generalized hypotonia, clenched hands, microphthalmia, large ear 7q11.22 | chr7:69718336-69783020 | Loss 64 kb | AUTS2 | Maternal |
| TM54-3 | M   | Non-syndromic ASD | None 8q24.3 | chr8:142,039,892-143,535,189 | Loss 1.49 Mb | PTP4A3, GPR20 | Paternal |
| TU29   | F   | ASD, ID | Facial dysmorphism, seizures 15q26.2 | chr15:95048523-96519797 | Gain 1.47 Mb | Reference genes: LOC440311, LINCO0924 | Maternal |
| TU23   | F   | ASD, ID, MCA | Microcephaly, brain abnormalities, facial dysmorphism, congenital heart defect, abnormal hearing test, congenital cataract 18q21.31-18q21.32 | chr18:55411917-56384192 | Gain 972 kb | ATP8B1, NEDD4L, MRRN122A, MALT1 | Paternal |
| RA39   | M   | ASD, ID | None 20p12.1 | chr20:14562027-14714457 | Loss 152 kb | MACROD2 | Paternal |
| TU4    | M   | Non-syndromic ASD | Short philtrum, 2 hair whirls, small eyes Xp22.31 | chrX:6449601-8141017 | Gain 1.69 Mb | HDHD1A, ST5, VCH, XCV2, XCV3A, PNP4 | Maternal |
| RA6    | M   | ASD, ID | None | chr15:4735698-155094135 | Gain 358 kb | TMLHE, SPRY3 | Maternal |

Table 2. Summary of all pathogenic CNVs and VOUS identified in this study. Abbreviations: VOUS, variant of uncertain clinical significance; ASD, autism spectrum disorder; ID, intellectual disability; DD, developmental delay; GDD, global developmental delay; MCA, multiple congenital anomalies; NA, not available.
and used various microarray platforms with different resolutions. A summary of microarray studies in patients with ASD is given in Supplementary Table S3. Of note, because the diagnostic yield of karyotype abnormalities in ASD has been shown to be lower than 3%7,23, it is not surprising that there were no statistically significant differences in the detected frequencies of pathogenic CNVs and VOUS between group 1 patients with normal karyotype and group 2 patients with the use of CMA as the first-tier test. Therefore, our findings support the recommendation of using CMA as the first-tier diagnostic test in patients with ASD of unknown cause.

Among pathogenic CNVs, the 16p13.11 microduplication was initially considered as a rare benign CNV38 however, we interpreted this duplication as pathogenic because several previous studies have supported the positive association between this duplication and a variety of neuropsychiatric disorders including ASD, unexplained ID, schizophrenia, epilepsy, and attention-deficit hyperactivity disorder (ADHD). This duplication region contains two strong candidate genes, NTAN1 and NDE1, which are both expressed in the brain and have been proposed as candidate genes for ASD and other neuropsychiatric disorders39–42. Most of the pathogenic, likely pathogenic CNVs and VOUS identified in this study were transmitted from a healthy parent (18 cases from 23 families, from which parental DNA samples were available). These inherited CNVs may suggest incomplete penetrance or some other factors interacting with the CNVs. Likewise, pathogenic CNVs in 1q21.1-1q21.2 (duplication)43–45, 15q13.2-15q13.3 (deletion)46,47, and 16p13.11 (duplication)40,41 were also found to be inherited in our study. The other possible theory would be the two-hit model, i.e. a second mutation not detected by CMA 48. Our findings further emphasize that parental array study alone may not be conclusive enough for CNV interpretation, the VOUS found in microarray studies need to be further analyzed before making a definite conclusion as to whether they are pathogenic or benign CNVs.

In the five de novo CNVs, one was a large deletion associated with the known 9q21.13 microdeletion syndrome, while the other four were duplications ranging from 127–220 kb in size. These duplications contained recently identified ASD-associated genes, DPP1049 and CHL149,50, and two genes that are highly expressed in the brain, but have not yet been linked to ASD, namely carnosine dipeptidase 1 (CNDP1) and serine incorporator 2 (SERINC2). Duplications of neither SERINC2 nor CNDP1 have been reported in the Thai CNV database51. The 1p35.2 duplication containing SERINC2 in particular is rare, and is not seen in the DGV database. Loss-of-function (LoF) variants (nonsense, frameshift and splice site variants) of these two genes were not observed in the 195 unrelated Thais from our in-house exome sequencing database (data not shown). In addition,
16 rare LoF variants of SERINC2 (MAF < 0.001) and 27 rare LoF variants of CNDP1 (MAF < 0.001) were found in the Exome Aggregation Consortium (ExAC) database. However, clinical phenotype data was not available for these individuals.

A de novo duplication of chromosome 18q22.3 included the ZNF407, CNDP2 and CNDP1 genes. Balanced translocation and point mutations in the ZNF407 gene were identified in one study of ID patients with autism32, but deletions/duplications of ZNF407 and CNDP2 genes have not been previously reported in patients with autism. The Carnosine dipeptidase 1 gene (CNDP1, OMIM 609064), also known as serum carnosinase (CN1), is involved in neuroprotective actions and neurotransmitter functions in the brain. It is mainly expressed in the cytosol of pyramidal neurons in the hippocampus and in large and small neurons of the temporal cortex in the human brain33. Alterations of serum carnosinase expression and activity have been shown to be associated with neurological diseases including multiple sclerosis and Parkinson’s disease34. Although carnosinase deficiency has been reported in patients with ID and DD35–37, the overexpression of CNDP1 from copy number gain as seen in our patient has never, to our knowledge, been previously reported. Additionally, carnosinase is an enzyme whose alterations is likely to be recessively inherited, so CNDP1 gene duplication may not affect the carnosinase function. A search of the DGV database found no CNVs which encompassed the exact same duplicated region, but we did find several smaller duplications encompassing CNDP1 in the general population. A search of the DECIPHER database did not reveal cases with deletion and duplication relatively similar in size to this duplication region. In addition, two DD cases (nsv581777 and nsv707115) with 178.39 and 279.02 kb duplications of the CNDP1 region were submitted to the ISCA/ClinGen database and interpreted as benign and uncertain-likely benign (Supplementary Figure 1a). Thus, these findings do not support CNDP1 as an ASD candidate gene.

A de novo 220 kb duplication of chromosome 1p35.2 encompassed the FABP3, SNRNP40, NKAIN1 and SERINC2 genes. Although FABP3 knockout mice by genetic and novelty seeking48 and one study found the NKAIN1 protein strongly expressed in the hippocampus and cerebellar granular cell layer39, no evidence for association between duplication of FABP3, SNRNP40, and NKAIN1 genes and neurodevelopmental disorders has been reported. Of particular interest is the serine incorporator 2 gene (SERINC2, OMIM 614549), also known as tumor differentially expressed 2-like (TDE2), which encodes a transmembrane protein that facilitates incorporation of serine into phosphatidylserine and sphingolipids50. The concentration of sphingolipids is highest in the brain, they play important roles in neural plasticity, signaling and axonal guidance51. Expression of Serinc2 mRNA was upregulated in the dentate gyrus of the hippocampus and the cerebellar Purkinje cell layer following kainate-induced seizures in rat52. In addition, overexpressed SERINC2 protein was reported in patients with developmental delay carrying deletions or mutations of the UPP2 or UPP3 genes, both of which are implicated in nonsense-mediated mRNA decay pathway (NMD)53. Furthermore, rare SERINC2 variants were significantly associated with alcohol dependence in subjects of European descent54. Transcript expressions of genes in the NKAIN1–SERINC2 genomic region were significantly associated with expressions of genes in the dopaminergic, serotonergic, cholinergic, GABAAergic, glutamatergic, histaminergic, endocannabinoid, metabolic, neuropeptide and opioid pathways55. Several studies have reported that alcohol dependence was associated with increased risk of neuropsychiatric disorders, including bipolar disorder and autism56–58. Autism and alcohol dependence have been shown to share some genetic factors according to earlier studies, where a higher incidence of alcoholism in the family members of ASD patients compared with the general population was observed. Also, the links between the autism susceptibility candidate 2 gene (AUTS2), a known ASD risk gene, in the regulation of alcohol consumption was reported59,60. These findings support the hypothesis that SERINC2 may be one of the dosage-sensitive genes and an increased SERINC2 protein may predispose to neurodevelopmental and neuropsychiatric disorders partly via abnormal NMD pathways and neurotransmitter or metabolic systems.

In the DECIPHER database, we identified two CNVs of slightly larger size encompassing SERINC2, a 377.2-kb microdeletion that has been reported in a patient with seizure (patient 289527 in Supplementary Figure 1b), and a 434.6-kb microduplication reported in a patient with ID and short attention span who also has Potocki-Shaffer syndrome. While the microdeletion was deemed possibly pathogenic, the microduplication was inherited from healthy parent, raising the possibility of incomplete penetrance. In addition, two larger duplications containing SERINC2 were seen in the ISCA/ClinGen database. These are a de novo 1.27-Mb duplication identified in a patient with global DD that was interpreted as pathogenic (nsv578518), and a 1.15-Mb VOUS identified in a patient with dysmorphic facial features (nssv1602204). Although the effects of other genes in these large duplication regions in the ISCA database could not be excluded, SERINC2 may contribute to neuropsychiatric disorders as noted in the supportive evidence discussed above. Because this gene have not been previously linked to autism, we proposed the SERINC2 gene as a novel ASD candidate gene. The supportive evidences of SERINC2 as a potential ASD candidate gene are summarized in Supplementary Figure 4.

Alpha-thalassemia (α-thalassemia) is the most common incidental finding of CMA in our Thai population. In Northern Thailand and Laos, the prevalence of α-thalassemia was estimated as between 30–40% including 3.6–10% of α-thalassemia 1 and 16.4–20% of α-thalassemia 2.69. Since the majority of alterations in α-globin genes are deletions, a CMA with enough resolution should be able to detect these mutations. In this study, CMA testing could detect α-thalassemia in 9 of 114 ASD patients (7.89%); 4.39% α-thalassemia 1 (5/114), 1.75% α-thalassemia 2 (2/114), and 0.88% Hb H disease (1/114). A comparison of our CMA, which was a high resolution SNP array, and traditional single-tube multiplex PCR results indicated that CMA could accurately detect HBA gene deletion. In previous studies, oligonucleotide microarrays with custom design probes were successfully used to detect both α-thalassemia and β-thalassemia70–72. Thus, we suggest that a tentative diagnosis of α-thalassemia should be considered in patients who underwent CMA testing because this is important for genetic counseling and prevention of α-thalassemia, particularly in populations where the disease is endemic. Of note, distal deletion of chromosome 16p (16pter–16p13.3) causes alpha-thalassemia-intellectual deficit syndrome linked to chromosome 16 (ATR-16 syndrome, MIM 141750). Patients with ATR-16 syndrome present with either
alpha-thalassemia trait or Hb H disease associated with mild to moderate ID, and with dysmorphic features in some cases. Deletion at 16p13.3 includes the loss of multiple genes including HBA1 and HBA2 resulting in alpha thalassemia phenotype, however the genes responsible for ID and other development abnormalities are not yet clearly identified. Some studies have suggested that deletion of SOX8 at chromosome 16p13.3 could contribute to ID in patients with ATR-16 syndrome because of its high expression in the brain75,76. However, our ASD patients, who have alpha thalassemia trait due to 16p13.3 deletions ranging from 3,924–149,072 bp, were unlikely to have ATR-16 syndrome because the 16p13.3 deletions in previously reported patients were much larger (>800 kb)74,76 and included many other genes including SOX8.

We also reported an atypical PWS case with ASD caused by a UPD of chromosome 15. Although, the results of the microarray and MS-PCR analyses were consistent with Prader-Willi syndrome (PWS), the clinical features of this patient were milder than in typical PWS patients. Various studies have found that PWS patients with UPDs of 15q11-q13 had higher IQs, milder behavior problems77,78, and more pronounced clinical features of ASD than PWS patients with deletion of 15q11-q1379. Our patient carried two AOH regions at 15q11.1–15q22.2 and 15q26.1-15q26.3. The B-allele frequency (BAF) plot indicated both uniparental isodisomy and heterodisomy regions on chromosome 15, demonstrating results of the recombination process in meiosis (Supplementary Figure 1). From the BAF plot, we can also deduce that the non-disjunction occurred in maternal meiosis II, followed by post-zygotic loss of the paternal chromosome 15 by trisomy rescue80. Due to the many genes located within these two large AOH regions, there is a possibility that in addition to the maternal UPD15 there exists the interplay between multiple genes that may contribute to the atypical phenotype of this patient. However, more investigations are required to further elucidate the underlying pathogenesis.

In conclusion, our study is the first CMA testing in a large cohort of 114 patients with ASD from Thailand, a country which has not been well represented in ASD studies. Our findings support the usefulness of CMA as a routine diagnostic test in patients with ASD of unknown cause. The CMA information can help not only medical management, but also in providing appropriate genetic counseling and evaluation of recurrence risk in families. In addition, we propose that the SERINC2 gene is a novel ASD candidate gene. However, because of the limitation of the number of studies related to SERINC2, further investigations are required to confirm the effect of SERINC2 in patients with ASD.

Methods

**Patients.** A total of 114 Thai patients with ASD of unknown cause were recruited in the study, 91 males and 23 females, with ages ranging from 1 to 18 years at the time of recruitment. They were divided into two cohorts. The first cohort (Group 1) was first evaluated for the common genetic causes of ASD and all patients, selected from a previous report7, had normal karyotype and negative DNA tests for Fragile X syndrome and MECP2 mutations (defined as “non-syndromic ASD”). Therefore, they represented non-syndromic ASD patients with the use of CMA as a second line test. Group 1 patients comprised 68 retrospective cases with non-syndromic ASD, who met the DSM-IV criteria for ASD including 61 non-syndromic ASD with ID patients (nonverbal IQ < 70) and 7 non-syndromic ASD without ID patients (nonverbal IQ ≥ 70). The second cohort (Group 2) included 46 prospective ASD and DD cases with a clinical diagnosis of ASD based on the Diagnostic and Statistical Manual of Mental Disorders, Fifth Edition (DSM-5) criteria. DD is defined as performance at least two standard deviations below the age-appropriate mean and is not quantifiable with IQ testing. Group 2 patients were referred for CMA testing at Ramathibodi Hospital, Bangkok according to the American College of Medical Genetics and Genomics (ACMG) recommendations for the use of CMA as the first-tier diagnostic test in patients with unexplained ASD and DD.

The study was approved by the Institutional Ethics Committee of Faculty of Medicine, Prince of Songkla University (REC 48/364-006-3) and Faculty of Medicine Ramathibodi Hospital, Mahidol University (ID 12-57-03). The methods were carried out in accordance with the approved guidelines and relevant regulations. Written informed consent was obtained from the parents of each ASD individual.

**Chromosomal Microarray Analysis.** CMA was performed with the Infinium CytoSNP-850K v1.1 Beadchip (Illumina, San Diego, California, USA), according to the manufacturer’s instructions. The array contains approximately 850,000 single nucleotide polymorphisms (SNPs) markers spanning the entire genome with an average probe spacing of 1.8 kb. The data were analyzed using BlueFuse Multi software v.4.3 and GenomeStudio Data Analysis Software v. 2011.1 based on the reference human genome (hg19/GRCh37). Microarrays were carried out in available parents of the patients with pathogenic CNVs and VUS to investigate the inheritance pattern in the families using either the Infinium CytoSNP-850K Beadchip (850,000 SNP markers) or HumanCytoSNP-12 DNA Analysis BeadChip v2.1 kit (~300,000 SNP markers) depending on the size of the interested CNVs.

**CNVs interpretation.** In this study, only detected CNV with at least 10-probe coverage were reported. The detected CNVs were classified into pathogenic, benign, or variant of uncertain clinical significance (VUS) according to established guidelines from the International Standard Cytogenomic Array (ISCA) and the ACMG54. The classification was based on the size of CNV, gene content, the inheritance pattern, and information in the medical literature and public databases. The Database of Genomic Variants (DGV) and the Thai CNV database (http://www.thaicnv.icbs.mahidol.ac.th/thaicnv/index.php)44, which contains CNVs from 3,017 Thai individuals, were used to exclude common structural variations in the Thai population, while the Online Mendelian Inheritance in Man (OMIM) database was used to identify disease-causing genes. The ISCA and Database of Chromosomal Imbalance and Phenotype in Humans using Ensembl Resources (DECIPHER) were used as reference for known microdeletion and microduplication syndromes. In addition, three ASD-related
Methylation-specific PCR (MS-PCR) for uniparental disomy detection. DNA samples were treated with sodium bisulfite, followed by SNRPN methylation specific PCR (MS-PCR). MS-PCR was performed with original primer sets designed by Kubota et al.37 and alternative primer sets designed by Hassan Askree et al.38. Untreated DNA was used as a control to ensure complete sodium bisulfate conversion of DNA, and amplified using primer set as previously described39. Subsequently, the PCR products were run on 2.5% agarose gel and visualized through ethidium bromide staining under a UV transilluminator.

Statistical analysis. All statistical comparisons were made by Chi-square test using the online GraphPad QuickCalcs software (https://graphpad.com/quickcalcs/contingency1.cfm). An alpha level of 0.05 was used for all statistical analyses.

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Acknowledgements
We would like to thank the patients and their families who participated in the study. We also thank Prof. Dr. Wasun Chantratita for providing the in-house Thai WES database. This work was supported by grants from the Faculty of Medicine, Prince of Songkla University (48/364-006-3) and the National Center for Genetic Engineering and Biotechnology (BT-B-01-MG-18-4814) (P.L.), the Development Potentials of Thai people with intellectual disabilities Project, Faculty of Medicine Ramathibodi Hospital, Mahidol University (ID 12-57-03) and the Crown Property Bureau Foundation (N.J.). T.T., D.W., and N.J. are recipients of the Research Career Development Awards from the Faculty of Medicine at Ramathibodi Hospital.

Author Contributions
A.H. performed the experiments, analyzed data and co-wrote the manuscript. W.T. performed the experiments. T.T., K.R., T.H., T.S., R.R., J.W., J.C., D.W. and N.R. performed the clinical diagnoses for samples recruitment. S.F. performed experiments for confirmation of α-thalassemia. P.L. and N.J. supervised the project, designed the study, obtained funding, analyzed data and co-wrote the manuscript. All authors read and approved the final manuscript.

Additional Information
Supplementary information accompanies this paper at https://doi.org/10.1038/s41598-017-12317-3.

Competing Interests: The authors declare that they have no competing interests.

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