Prader-Willi syndrome (PWS) is a well described clinical dysmorphic syndrome but due to some overlap of features with both normal obese individuals and those with other developmental handicaps (1-5), a definitive diagnosis on clinical grounds alone is uncertain and requires DNA testing. There are various molecular genetic tests available and DNA testing is approximately (~)99% accurate for the diagnosis of PWS—this ranks as one of the best percentage accuracy of testing for any syndromic genetic condition. Thus, while the actual diagnosis is straightforward on the whole, the dilemma arises because of the complex genetic mechanisms at the basis of PWS, which need to be elucidated. To establish the molecular mechanism with a complete work up, involves at least 2 tests. Here we discuss the commonly used tests currently available and suggest a cost—effective approach to diagnostic testing.

Keywords: Cytogenetics; chromosome microarray; methylation

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Introduction

Prader-Willi syndrome (PWS) is a well described clinical dysmorphic syndrome but due to some overlap of features with both normal obese individuals and those with other developmental handicaps (1-5), a definitive diagnosis on clinical grounds alone is uncertain and requires DNA testing. There are various molecular genetic tests available and DNA testing is approximately (~)99% accurate for the diagnosis of PWS—this ranks as one of the best percentage accuracy of testing for any syndromic genetic condition. Thus, while the actual diagnosis is straightforward on the whole, the dilemma arises due to the complexity of establishing the molecular mechanism. It is necessary to establish the mechanism as this will provide information on the possible clinical features (4), the prognosis (6-8) and the recurrence risk (9,10). While there are various molecular tests available which can establish the mechanism, there is no set standard algorithm of testing; different laboratories use different approaches and often their own set pattern of testing, making the interpretation of test results difficult for clinicians. Here, the aim is to lay out the various possible tests currently in use and show the value of each one individually and in combination, to provide a useful guide for clinicians in understanding the genetic scenario of PWS diagnostic testing.

This discussion does not include a clinical appraisal of PWS, first described in 1956 by Prader et al. (11), which has been given in many previous publications since (1,4,12-15). The basic defect is in the malfunctioning of the hypothalamic nuclei controlling appetite. PWS occurs in approximately one in 20,000 live births (16) and occurs equally in males and females and in all ethnic groups. Angelman syndrome with a different phenotype, paternal imprinting, similar genetic mechanisms but extended to include a known gene (UBE3A) (17) is not considered in detail, but mentioned where it impinges on the interpretation of genetic testing for PWS.

The genetic abnormality in PWS

The genetic region for PWS on chromosome 15 was first
recognised through chromosomal rearrangements involving the centromeric region of chromosome 15 (18). Since then, the critical region for PWS has been established to lie within chromosome 15(q11-13) [OMIM No. 176270] a region spanning 4–6 Mb (Figure 1). A number of genes have been mapped within this region, the critical one for PWS is SNURF/SNRPN gene, which contains 10 exons, many snoRNAs, the imprinting centre (IC) and the open reading frame (ORF) of the gene. Of these, exon 1 is critical for imprinting (10,19-20) and SNORD 116 RNA is important for many of the clinical features of PWS (21,22) (Figure 1). PWS is maternally imprinted and is caused by loss of paternal alleles within this region (9,23). However, as yet no abnormal gene product has been identified which could be the cause of PWS. SNRPN is a small nuclear ribonucleoprotein involved in alternate mRNA splicing.

The critical region for Angelman Syndrome (AS) lies within this same 4–6 Mb region, but AS is caused by loss of maternal alleles (17).

The genetic mechanisms leading to PWS are 3 fold—either there is a deletion of the critical region (~75% of patients), uniparental disomy (UPD) (~25%) or an imprinting centre (IC) defect (~1% of PWS) (9,19,20,24) (Figure 2). These mechanisms can be arrived at by occurring de novo or as a result of a structural abnormality, such as a translocation, ring formation, isochromosome, or inversion, involving chromosomes 15 alone or with some other chromosome. A rearrangement occurs in ~5% of individuals with PWS (25). These structural rearrangements may be balanced or unbalanced, the commonest being Robertsonian translocations (14;15) and (15;15) (25-30).

Deletions
The PWS chromosome critical region spans 4–6 Mb and deletions within this region may be large, small or microdeletions. Large deletions of approximately 4–6 Mb have two major proximal breakpoints but a consistent single distal breakpoint (4,31). Class I deletions extend from D15S541 to D15S12 (BP1 to BP3) and class II from D15S543 to D15S12 (BP2 to BP3)—both take out the IC. (Figure 1). These deletions are considered collectively as the common deletion; there are subtle phenotypic difference between them (32); they are present in approximately equal proportions, and are the mechanism of PWS in ~75% of patients; there is no report of recurrence (4,19). Small deletions are those smaller than 4 Mb; their effect is similar
to a large deletion and there is also no report of recurrence. Microdeletions are within the SNRPN gene and span from 6–250 kb DNA, with exon 1 deleted (20,33). These are sometimes referred to as IC mutations (or IC defects) (see below) and have a potentially high (50%) recurrence risk (10,34).

**UPD**

The chromosome count is normal, with normal appearing chromosomes 15—band for band—but both have come from the mother—there is no paternal input. UPD (mat) in PWS was first described in 1989 (24) and while initially thought to be rare, accounts for ~25% of individuals with PWS. In UPDmat, the commonest mechanism is frequently referred to as a trisomy rescue—there are two abnormal events, the first is meiotic, resulting in a zygote with trisomy 15 and the second is somatic after fertilisation with loss of the paternal chromosome 15. Heterodisomy occurs when the 2 chromosomes 15 in the trisomy 15 zygote are different; isodisomy is the term used when they are the same. Advanced maternal age is correlated with UPD as the first error is a maternal non disjunction event, leading to maternal meiotic trisomy 15 (35). There are other rarer chromosome breakage and segregation abnormalities, which can result in UPDmat (30).

**Imprinting centre defect (ICD)**

Essentially ICD (also referred to as IC mutations) are microdeletions within SNRPN, involving exon 1. The size of these deletions varies only a little, from exon 1–10. DNA testing is the only way to detect ICD, and this is usually performed in a specialised referral laboratory (20,33). Fortunately, ICD occur rarely (~1% of individuals with PWS), as this testing may take some time and may involve another sample from the patient and one from each parent. If the mechanism of the common 4–6 Mb deletion or UPD is established, it is unnecessary to look for an ICD; otherwise, it is necessary, due to the high recurrence risk (50%), if it is found to be inherited. Testing for an IC deletion can be done by sequence analysis at the smallest region of overlap (SRO).
for the PWS IC region (~4.3 kb) (33,36).

Absence of the paternally derived SNORD116 gene cluster has been described as a rare cause of PWS with multiple clinical features as seen in typical PWS (37). Most (85%) of the imprinting centre defects are due to epimutations (i.e., two parental alleles but only maternal DNA methylation pattern), with small recurrence risk. The remaining ~15% of PWS with imprinting centre defects are found to have very small deletions in the IC region (7.5~>100 kb) with recurrence risk as high as 50% if the IC deletion is inherited from an unaffected father (10,23,24).

The dilemma for diagnostic genetic laboratories

There are 4 issues to consider:

(I) To separate out the few patients with PWS from the great majority of laboratory referrals falling within the differential diagnosis, at birth and later (1,3,38,39);

(II) Having diagnosed PWS, to separate out those few individuals with a high recurrence risk (an ICD) from the much larger number with a very low recurrence risk (deletions and UPDmat);

(III) To make the testing as “user-friendly” as possible—it is notoriously difficult to obtain blood, or other tissue, from the patient, and family is not always available;

(IV) To keep the cost of testing to a minimum.

Tests commonly utilised at present

Cytogenetics

Cytogenetics after setting up tissue culture (usually from a blood sample) chromosomes are released from their cells (hypotonic and colchicine treatment), spread on a slide (cytogenetic harvest), and banding techniques applied which can identify each chromosome accurately band for band to a level of resolution of 5–10 Mb. This is a useful test to check for structural rearrangements (e.g., Robertsonian translocations or small supernumerary chromosome 15 which are associated with ~5% of UPD) (30,36,40); it is not useful for the detection of deletions or UPD (40).

Fluorescence in-situ hybridisation (FISH)

FISH utilises a fluorescence microscope to visualise with a bright signal whether a probe hybridises to an homologous sequence or not—i.e., deleted or not deleted (Figure 3). Several probes from within the critical region can be used for FISH studies (e.g., SNRPN & D15S10). FISH, with the appropriate probes, is a useful way to differentiate type 1 from type 2 deletions. These probes can be applied to chromosome spreads prepared after cell culture or directly to smears, which are whole cells on a slide without culture. FISH can be performed on cells in the suspension remaining from the cytogenetic harvest, retained in fixative at ~20 °C (41). Thus, slides can be made later, using the same suspension, which can be useful for checking on some other result e.g., microarray. FISH on chromosome spreads effectively can characterise a balanced structural abnormality and is the test of choice for this purpose. FISH cannot be used to pick up UPD and will not distinguish between PWS and AS deletions.

Methylation polymerase chain reaction (PCR)

Methylation PCR can diagnose PWS unequivocally when the abnormal pattern is present, and can also exclude AS (Figure 4). Methylation does not define the PWS molecular mechanism, as it is abnormal in all 3 scenarios—the common large and small deletions, UPD and IC defects (42,43). When an abnormal methylation result is shown, further testing is required to indicate the mechanism. This further testing is commonly done with FISH—a single probe will detect a deletion; if the FISH result is normal i.e., not deleted, DNA testing is required to show UPD. If UPD is not shown, an imprinting defect is suspected,
and further special testing is necessary if the IC defect is to be characterised. At present >95% of IC defects are microdeletions within SNRPN (33). Rare patients have been reported who have PWS microdeletion by chromosome rearrangement (28,29).

Comparative genomic hybridisation (CGH) microarray

CGH microarray compares the DNA of a patient with a normal control. It can be either a CGH array (detects copy number change only) or CGH + single nucleotide polymorphism (SNP) array which can detect both copy number change as well as single nucleotide polymorphism change. Both can detect deletions of all sizes and unbalanced chromosome rearrangements (Figure 5). The whole genome is visible with microarrays, not just the PWS region, so that other abnormalities elsewhere in the genome can be detected. The precise deletion size and its corresponding genomic content may be important for genotype-phenotype correlations. CMA combined with SNPs (CGH + SNP array) can detect approximately 75% of UPDs by demonstrating long (>13.5 Mb) contiguous stretches of homozygosity limited to the chromosome 15 s (44). In these cases DNA polymorphism analysis is probably not necessary to diagnose UPD. One of the limitations of CGH + SNP is that it cannot detect all cases of uniparental heterodisomy UPD or a balanced chromosome rearrangement (5% of patients) (45). Furthermore, interpretation of the microarray result may be inaccurate if the patient has a homozygous mutation or mosaicism (46), both quite rare in PWS.

Methylation sensitive multiplex ligation-dependent probe amplification (MS-MLPA)

Each MS-MLPA reaction generates two samples that require analysis by capillary electrophoresis:

One is an undigested sample to check for copy number change and the other is a digested sample (with HhaI endonuclease enzyme) for methylation detection. MS-MLPA can detect copy number changes as well as determining methylation status of the 15q11 region (47). It can also pick up small imprinting centre (IC) deletions and microdeletions within the SNORD116 cluster (Figure 6). MS-MLPA will detect UPD, but cannot definitively distinguish UPD from imprinting centre mutation.

![Figure 4](image_url) Methylation-PCR DNA was modified by bisulfite treatment and the SNRPN exon 1 region was amplified by PCR. Normal control has both the maternal (313 bp) and paternal (221 bp) bands. Only the maternal band is visible in a Prader-Willi patient and the paternal band is present in an Angelman patient. PCR, polymerase chain reaction.
Figure 5 CGH chromosome microarray with type 1 deletion [between BP1–BP3 (GRCh37 chr15:22,749,354-28,438,266)], the CGH data showing Log2 ratio “−1” between chromosome 15q11. 2q13. 1 regions indicating a heterozygous deletion. CGH + SNP chromosome microarray with type 2 deletion [between BP2-BP3 (GRCh37 chr15:23,619,912-28,438,266)], the CGH data showing Log2 ratio “−1” between chromosome 15q11. 2q13. 1 region indicating a heterozygous deletion. The SNP data also showed loss of one allele [displaying only the 0-uncut (AA) & 1-uncut allele (AB) track]. (A) CGH microarray-Type1 deletion; (B) CGH + SNP Microarray-Type2 deletion.
Figure 6 MS-MLPA showing normal control, PWS large common deletion, IC deletion and UPD.
RNA

The measurement of the expression of the SNRPN gene is another approach and considered by some to be more important than the actual presence of an abnormality (48). This is a complicated growing area as there are many RNA transcripts (with no apparent protein coding function) apart from the classical messenger RNA. There are 3 non coding RNA (ncRNA) of importance in the PWS region (HB13, 52 and 85) (Figure 1)—these may not be translated because the RNA binds to other molecules, or is methylated. Starting with an RNA test (not shown), lack of gene expression will diagnose PWS, but there is still the need to establish the mechanism and a cytogenetic analysis is also required to check for a balanced rearrangement. RNA extraction may be unreliable and extracted RNA unstable, so that this approach is not commonly used for diagnosis.

Testing algorithms

A comprehensive, definitive diagnosis requires a minimum of two tests, and depending on the choice of tests, these same two tests could exclude a balanced rearrangement. These two tests could be performed from one given specimen, or from 2 specimens taken at different points in time. If the specimen was sufficiently large, e.g., 10–15 mLs, then it could be split—part for a cytogenetic karyotype analysis or FISH, looking for a balanced chromosome rearrangement and the other part for DNA studies.

Starting with the DNA methylation test (from DNA extraction) a diagnosis will be made immediately, and AS excluded, but further testing is required for the mechanism. Frequently, FISH is the next test, for the common 4–6 Mb deletion as the mechanism, (from cells on a microscope slide, either from a smear or cell culture); this will cover 75% of patients. If the FISH test is normal, then proceed to DNA studies for UPD; this entails going back to the original specimen, or obtaining another specimen, or using the cell suspension remaining from the cytogenetic harvest and extracting DNA; then also obtaining blood from the parents (4,49).

Starting with a microarray, a deletion will be detected immediately but will not distinguish PWS from AS. A CGH + SNP microarray can detect UPD by loss of heterozygosity, but cannot determine the parent of origin for UPD unless a trio-analysis of the proband and both parents are performed simultaneously. However, if the UPD15 has resulted from a maternal uniparental heterodisomy (trisomy rescue) then, a trio-analysis would not be useful. Microsatellite analysis in the patient and parents are still required. However, a CGH array has an advantage in that it will detect abnormality across the whole genome.

Starting with MS-MLPA, a definitive diagnosis can be achieved in one test for ~75% of patients (Figure 6). The diagnosis plus deletion is achieved without the need for parental DNA and can distinguish AS from PWS, detecting all types of deletion (4,15). The limitation is that cases with UPD require another molecular test (either CGH + SNP array to detect long contiguous stretches of homozygosity, or DNA polymorphism studies by microsatellite analysis). The parents are still required to check for UPD. Thus, MS-MLPA cannot discriminate between UPD and IC epimutation. MS-MLPA assay does not detect copy number changes across the entire genome and it also cannot detect balanced rearrangements. Due to the rarity of ICD and balanced rearrangements in PWS, some centres consider it sufficient to diagnose PWS and establish deletion as the mechanism (50).

Cost

The cost of these various tests varies from one laboratory or genetic service to another. Some will charge for a complete diagnosis, others charge for each step of the process (e.g., DNA extraction, a cytogenetic harvest, slide making, cytogenetic analysis) so that a useful comparison cannot be made. The less tests required, the more cost effective the diagnosis becomes. Retesting after an incomplete diagnosis is costly (51). As a baseline for comparison (52), in our genetic service in Sydney, Australia, the cost of a cytogenetic karyotype is $A365.00; a CMA is $A365.00; FISH test is $A175.00; methylation test is $A300.00 and MS-MLPA is $A300.00. A charge can be made for one or the other of cytogenetic karyotype or CMA but not both, as the CMA is classified as a molecular karyotype. The patient may not need to pay for all this cost, as the Medicare scheme in Australia will cover a percentage of the cost (but not for the methylation or MS-MLPA) and some families have private health care cover. However, apart from the testing itself, there is considerable paperwork involved in tracking the
case for the laboratory scientist, and a secretarial cost should also be added.

Thus, a comprehensive laboratory diagnosis remains a dilemma; in Figure 7, we present a pathway which will definitely diagnose and elicit the molecular mechanism of PWS in a cost-effective and timely manner, with MS-MLPA as the first test. This will be valid for the majority of patients (75% deletion). The compromise is that if UPD studies are to be proven and not just assumed when no deletion is found, and a balanced rearrangement is also to be excluded, further testing is necessary and may involve another specimen and the parents.

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Footnote

Conflicts of Interest: The authors have no conflicts of interest to declare.

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