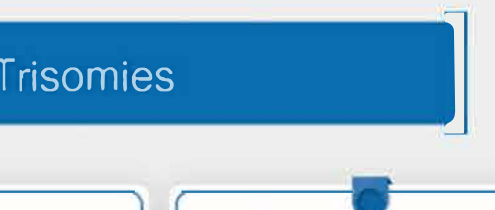


標準版

Standard Panel

為了使服務達到國際標準³，我們建議媽媽選擇
敏兒安^{safe}21 express 標準版，其所報的14項常見異常
有較多數據支持。

safe^{safe}21 express Standard Panel screens for 14 validated conditions. In alignment with international guidelines³, this panel is the recommended choice for most mothers.



測試項目包括 Testing items include

3 項

染色體三體症
Trisomies

7 項

微缺失症候群
Microdeletion Syndromes

4 項

性染色體相關疾病
Sex Chromosome Aneuploidies

染色體三體症 Trisomies

T21唐氏綜合症
T21 Down Syndrome

T18愛德華氏綜合症
T18 Edwards Syndrome

T13巴陶氏綜合症
T13 Patau Syndrome

微缺失症候群 Microdeletion Syndromes

1p36缺失綜合症
1p36 Deletion Syndrome

2q33.1缺失綜合症
2q33.1 Deletion Syndrome

15q11.2缺失綜合症/
天使綜合症
15q11.2 Deletion/
Angelman Syndrome

5p缺失綜合症/
貓哭綜合症
5p Deletion/
Cat-du-chat Syndrome

22q11.2缺失綜合症/
迪喬治綜合症
22q11.2 Deletion/
DiGeorge Syndrome

8q24.1缺失綜合症/
毛髮-鼻-指骨綜合症
8q24.1 Deletion/
Langer-Giedion Syndrome

15q11.2 缺失綜合症/
普瑞德威利綜合症/
小胖威利綜合症
15q11.2 Deletion/
Prader-Willi Syndrome

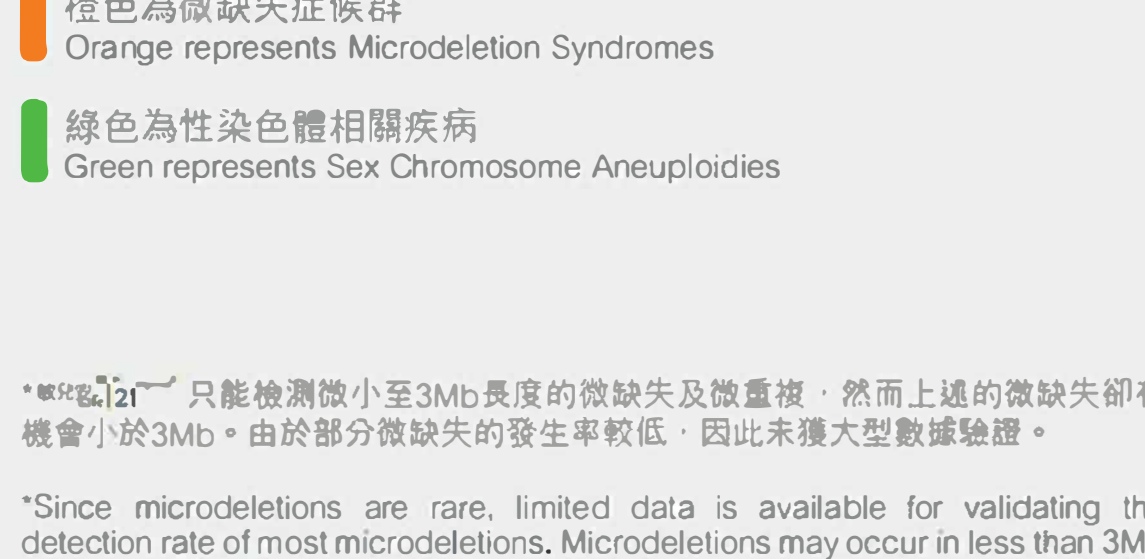
性染色體相關疾病 Sex Chromosome Aneuploidies

X0 X染色體單體症
(特納綜合症)
X0 Monosomy X
(Turner Syndrome)

XYY 三體綜合症
(XYY 超雄綜合症/
雅各氏綜合症)
XYY Syndrome
(Jacob's Syndrome)

XXY 柯林菲特氏綜合症
XXY Klinefelter Syndrome

XXX 三體綜合症
(XXX 超雌綜合症)
XXX Triple X Syndrome



藍色為染色體三體症
Blue represents Trisomies

橙色為微缺失症候群
Orange represents Microdeletion Syndromes

綠色為性染色體相關疾病
Green represents Sex Chromosome Aneuploidies

*敏兒安^{safe}21[™] 只能檢測微小至3Mb長度的微缺失及微重複，然而上述的微缺失卻有機會小於3Mb。由於部分微缺失的發生率較低，因此未獲大型數據驗證。

*Since microdeletions are rare, limited data is available for validating the detection rate of most microdeletions. Microdeletions may occur in less than 3Mb in size, safe^{safe}21[™] only searches for microdeletions with a minimum size of 3Mb.

進階版 Advanced Panel

敏兒安^{safe}21^{express} 進階版可以全面檢測到23對染色體相關疾病，包括染色體三體症及目前已在國際數據庫(OMIM, Decipher和Orphanet)中記錄的105項微小至3Mb的微缺失或微重複^{4, 5, 6, 7}。

safe21^{express} Advanced Panel screens for chromosomal aneuploidies of all 23 pairs of chromosomes, including trisomies and the 105 microdeletions/microduplications with a minimum size of 3Mb that has been recorded on the international databases: OMIM, Decipher and Orphanet^{4, 5, 6, 7}.

敏兒安^{safe}21^{express}

測試項目包括 Testing items include

22 項

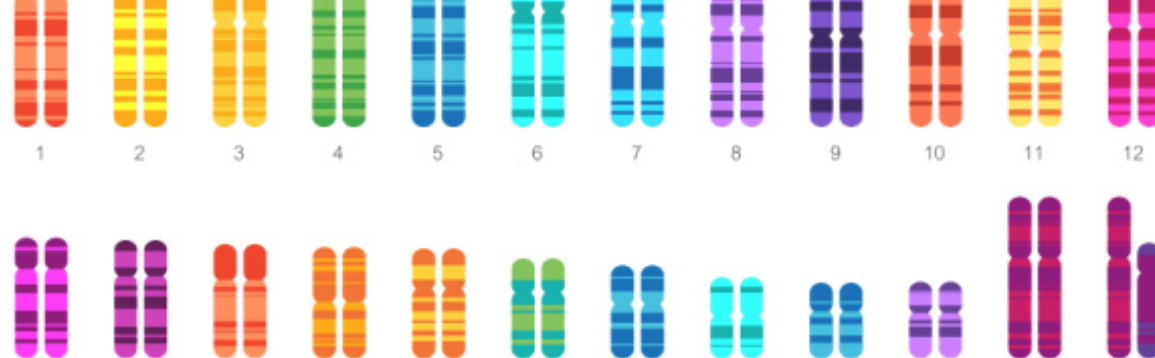
染色體三體症
Trisomies

105 項

微缺失或
微重複症候群
Microdeletion/
Microduplication
Syndromes

4 項

性染色體相關疾病
Sex Chromosome
Aneuploidies



22 項染色體三體症
22 Trisomies

XO
XYY
XXY
XXX

4 項性染色體相關疾病
4 Sex Chromosome
Aneuploidies

*敏兒安^{safe}21^{express} 只能檢測微小至3Mb長度的微缺失及微重複，然而上述的微缺失都有機會小於3Mb。由於部分微缺失的發生率較低，因此未獲大型數據驗證。

*由於進階版的檢測項目罕見及複雜，未獲大型數據驗證，因此準確率會降低。

*Since microdeletions are rare, limited data is available for validating the detection rate of most microdeletions. Microdeletions may occur in less than 3Mb in size, safe21^{express} only searches for microdeletions with a minimum size of 3Mb.

*Since advanced findings are rare and complex, insufficient data for validation may lower the accuracy.