

報告編號(No.): HTF24500978M03

報告日期(DATE): 2024/06/14 頁數(PAGE): 1 of 4

Test Report

集泉塑膠工業股份有限公司(LIVING FOUNTAIN PLASTIC INDUSTRIAL CO., LTD.) 台中市霧峰區民生路198巷31號(NO.31, LN. 198, MINSHENG RD., WUFENG DIST., TAICHUNG CITY 41346 TAIWAN)

以下測試樣品係由申請廠商所提供及確認 (The following sample(s) was/were submitted and identified by/on behalf of the applicant as): 送樣廠商(Sample Submitted By) : 集泉塑膠工業股份有限公司 (LIVING FOUNTAIN PLASTIC INDUSTRIAL CO., LTD.) 樣品名稱(Sample Description) : CPET PLATE (CPET潔晶餐盤)

樣品名稱(Sample Description) 樣品材質(Sample Material) 原產國(Country Of Origin)	:	CPET PLATE(CPET ※ 品資盤) CPET TAIWAN
 收件日(Sample Receiving Date) 測試期間(Testing Period)	:	2024/05/28 2024/05/28 to 2024/06/07

測試需求 (Test Requested)

- (1) 客戶指定依據美國聯邦法規之藥物暨食品管理 (FDA) 21 CFR 177.1630 (f) 進行測試。測試項目請參閱測試結果表格。 (As specified by client, the sample(s) was/were tested according to U.S. Food and Drug Administration (FDA) 21 CFR 177.1630 (f). Please refer to the result table(s) for the testing item(s).)
- (2) 客戶指定依據美國聯邦法規之藥物暨食品管理 (FDA) 21 CFR 177.1630 (g) 進行測試。測試項目請參閱測試結果表格。 (As specified by client, the sample(s) was/were tested according to U.S. Food and Drug Administration (FDA) 21 CFR 177.1630 (g). Please refer to the result table(s) for the testing item(s).)
- (3) 客戶指定依據美國聯邦法規之藥物暨食品管理 (FDA) 21 CFR 177.1630 (h) 進行測試。測試項目請參閱測試結果表格。 (As specified by client, the sample(s) was/were tested according to U.S. Food and Drug Administration (FDA) 21 CFR 177.1630 (h). Please refer to the result table(s) for the testing item(s).)

測試結果 (Test Results) : 請見下一頁。(Please refer to next page(s).)



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測試部位敘述(Test Part Description)

No.1 : 白色塑膠盤 (WHITE PLASTIC PLATE)

測試結果(Test Results)

(1) 通過(PAS					過(PASS)
測試項目 (Test Item(s))	測試方法 (Method)	單位 (Unit)	RL	結果 (Result)	限值 (Limit)
	(includy)	(onn)		No.1	()
氯仿可萃取物 (水, 250°F, 2小時) / Net chloroform-soluble extractives (D.I. Water, 250°F, 2 h)	參考美國FDA 21 CFR 177.1630 (f). / With reference to US FDA 21 CFR 177.1630 (f).	mg/in²	0.1	n.d.	0.5
氯仿可萃取物 (正庚烷, 150°F, 2小時) / Net chloroform-soluble extractives (n-Heptane, 150°F, 2 h)		mg/in²	0.1	n.d.	0.5

(2)

(=)				2	
測試項目 (Test Item(s))	測試方法 (Method)	單位 (Unit)	RL	結果 (Result) No.1	限值 (Limit)
氯仿可萃取物 (水, 250°F, 2小時) / Net chloroform-soluble extractives (D.I. Water, 250°F, 2 h)		mg/in²	0.1	n.d.	0.5
氯仿可萃取物 (50% 乙醇, 120°F, 24小時) / Net chloroform-soluble extractives (50% Alcohol, 120°F, 24 h)	參考美國FDA 21 CFR 177.1630 (g). / With reference to US FDA 21 CFR 177.1630 (g).	mg/in²	0.1	n.d.	0.5
氯仿可萃取物 (正庚烷, 150°F, 2小時) / Net chloroform-soluble extractives (n-Heptane, 150°F, 2 h)		mg/in²	0.1	n.d.	0.5

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通過(PASS)



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(3)	
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通過(PASS)

測試項目 (Test Item(s))	測試方法 (Method)	單位 (Unit)	RL	結果 (Result) No.1	限值 (Limit)
氯仿可萃取物 (水, 250°F, 2小時) / Net chloroform-soluble extractives (D.I. Water, 250°F, 2 h)	參考美國FDA 21 CFR 177.1630 (h). / With reference to US FDA 21 CFR 177.1630 (h).	mg/in²	0.01	n.d.	0.02
氯仿可萃取物 (正庚烷, 150°F, 2小時) / Net chloroform-soluble extractives (n-Heptane, 150°F, 2 h)		mg/in²	0.01	n.d.	0.02

備註(Note):

- 1. RL = Reporting Limit (報告極限值)
- 2. n.d. = Not Detected (未檢出) = Less than (小於) RL
- 3. 本報告不得分離或擷錄使用。(The report is invalid if it is partly reproduced or used.)
- 本實驗室之報告符合性聲明依ILAC-G8:09/2019簡單允收之二分法判定規則(w=0 · AL=TL)做為測試結果符合性聲明 判定之判定依據。(The decision rule of the statements of conformity is following the ILAC G8:09/2019 by using the simple acceptance decision rule.)
- 5. 本報告為 HTF24500978 之加發報告,報告加發日期 2024年06月14日。 (This is the additional test report of HTF24500978. The additional test report is issued on 2024/06/14.)

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> * 照片中如有箭頭標示,則表示為實際檢測之樣品/部位。 * (The tested sample / part is marked by an arrow if it's shown on the photo.)

No.1





** 報告結尾 (End of Report) **

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