



US FDA CDER DMF No.: 038447 US FDA CBER DMF No.: 29657 **Acute Systemic Toxicity Test** 

DASEA Ultramedia® CIK Cell Serum-Free Medium is speially designed to promote high-efficiency proliferation of CIK cells in vitro. DASEA Ultramedia® CIK contains no animal-derived components or heterologous growth factors which enhances experimental reproducibility by avoiding batch-to-batch variations caused by

\* For Research Use Only



High Proliferation Efficiency



High Positive Rate



**GMP** Manufacturing

### **ADVANTAGES**

### Safe and reliable raw material management



- Thousand-square-meter storage space for raw materials.
- Strictly monitor temperature and humidity of the warehouse.



### **Optimal batch-to-batch** stability

• Compliance with raw material release criteria.

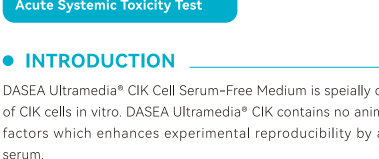
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STERILR A

DASEA Ultramedia® CIK细胞无血清培养基

本产品不含血清和动物源成分,提高实验可重复性 支持相关因子介导的体外CIK 的无血清扩增培养。 产品货号: RGL0040

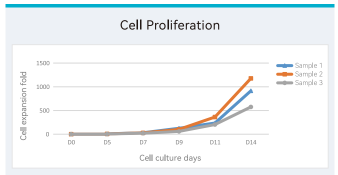
 Strictly define and enforce production process control parameters.



# QUALITY STANDARD

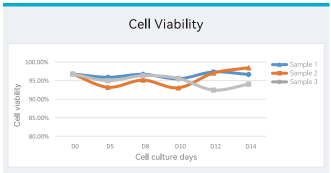
Quality Inspection Item	Quality Inspection Standards
Appearance	Orange-red clear liquid
рН	7.2 ± 0.2
Osmolality	280-320 mOsm/kg
Endotoxin	<0.5 EU/mL
Sterility Test	Negative
Mycoplasma Test	Negative

## PERFORMANCE DATA



D0: Total 10ml blood was used to collect CIK cells. Initial cell culture volume is 5mL and initial cell count is 7.5 x 10<sup>6</sup>.

D14: Cell number were increased by 500–1000 times. 3.7 billion to 7.5 billion cells were harvested, with 1.2–2L DASEA Ultramedia $^{\circ}$  CIK medium usage.



The cell viability kept high during the high density cell culture process.

