同意書

編號 NUM

Date

AS-IRB02- 101043

AS-IRB02- 10104.

日期 0402'2012

中央研究院醫學研究倫理委員會 研究計畫審查通過證明

Certificate of Approval

IRB02 修訂日期 Revised Date: 0812'2010

申請案編號 Application No:

AS-IRB01- 12029 (12029)

研發癌症標記胜肽應用於腫瘤之標靶藥物傳輸

計畫名稱 Project title:

有效期限 Due Date:

Development of Cancer Cell-Specific Peptide for Tumor Targeted

Drug Delivery

申請人 Project Investigator:

吳漢忠 Wu, Han-Chung

合作機構 Collaborating Institute(s): 無

執行期間 Implement Duration: From <u>2012 / 07 / 01 / to 2015 / 06 / 30 /</u>

(yyyy / mm/ dd) (yyyy / mm/ dd)

From 2012

From <u>2012</u> / <u>07</u> / <u>01</u> / to <u>2015</u> / <u>06</u> / <u>30</u> /

(yyyy / mm/ dd) (yyyy / mm/ dd)

進度或成果報告 Progress Report:

1. 請於 <u>2013 / 06 / 30 /(yyyy / mm/ dd)</u>前繳交進度或成果報告至本委員會追蹤審查。
The progress report should be submitted to the IRB at Academia Sinica before <u>2013 / 06 / 30 / (yyyy/ mm/ dd)</u> for continuing review.

- 2. 如向衛生署定期提出進度或成果報告,需將完整報告內容副知本委員會。
 If Progress reports are submitted to the Department of Health, a copy should be submitted to the IRB at Academia Sinica.
- 3. 如受試者有不良反應,應即時向主管機關報告並副知本委員會。 All the adverse events are required to report to the official institute and IRB of Academia Sinica promptly.

附註意見 Additional Opinion:

若需與受試者溝通(如與受試者通信、致贈禮物等),與受試者溝通之文件、信件或禮物,須經本委員會審核通過後方可進行。

Any materials (correspondence, brochure, compensation, etc.) provided to subjects, must be approved by the IRB of Academia Sinica.

茲 證明上項醫學研究計畫,包括其受試者之說明同意書,已經本委員會之審核,並同意此計畫書之內容符合本委員會所訂定之醫學研究倫理標準。

The above named study proposal has been reviewed, along with the informed consent, by the committee and found to conform with the guidelines set forth by this committee.

主任委員:

日期:

Chairman

Date

中央研究院醫學研究倫理委員會

Human Subject Research Ethics Committee / IRB Academia Sinica