

## Modified griffithsin tandemers for enhanced activity and reduced viral aggregation

### Summary (1024-character limit)

NCI seeks partners to commercialize Griffithsin and Griffithsin tandemers as therapeutics for HIV infections that are resistant to native GRFT, specifically, additional studies on stability, toxicity, immunogenicity, and large-scale production.

### NIH Reference Number

E-034-2013

### Product Type

- Therapeutics

### Keywords

- mGRFT tandemers
- antiviral

### Collaboration Opportunity

This invention is available for licensing.

### Contact

- John D. Hewes  
NCI - National Cancer Institute

240-276-5515

[John.Hewes@nih.gov](mailto:John.Hewes@nih.gov)

### Description of Technology

Griffithsin (GRFT) is a lectin with potent antiviral properties that is capable of preventing and treating infections caused by a number of enveloped viruses (including HIV, SARS, HCV, HSV, and Japanese encephalitis) and is currently in clinical development as an anti-HIV microbicide. In addition to its broad antiviral activity, GRFT is stable at high temperature and at a broad pH range, displays low toxicity and immunogenicity, and is amenable to large-scale manufacturing. Native GRFT is a domain-swapped homodimer that binds to viral envelope glycoproteins and has displayed mid-picomolar activity in cell-based anti-HIV assays.

Researchers at [NCI's Molecular Targets Lab](#) developed synthetic proteins that comprise two (or more) obligate monomers ("mGRFT") joined by an amino acid linker to form tandemers ("mGRFT tandemers"). Each obligate monomer is generated by the addition of Gly-Ser residues in the hinge region of wild-type GRFT. Two or more obligate monomers are joined by an amino acid linker to form the mGRFT tandemers.

The properties of the mGRFT tandemers can be modulated by the length of the amino acid linker and the number of obligate monomers co-joined. mGRFT tandemers exhibit more potent anti-viral properties when compared against native GRFT and are equipotent against viruses that are both sensitive and resistant to native GRFT. As such, potential uses of the invention tandemers include topical and intravenous therapy to treat HIV infection, particularly to treat HIV infections that are resistant to native GRFT.

### Potential Commercial Applications

- Broad-spectrum antiviral agent similar to wild type GRFT
- Potential activity against SARS CoV, MERS, Ebola, HCV and influenza

### Competitive Advantages

- Broad antiviral activity and stable at high temperature and at a broad pH range
- Displays low toxicity and immunogenicity

### Inventor(s)

[Barry R. OKeefe \(NCI\)](#), [Alexander Wlodawer \(NCI\)](#), [Tinoush Moulaei \(NCI\)](#)

### Development Stage

- Pre-clinical (in vivo)

### Publications

Moulaei T et al. Griffithsin tandemers: flexible and potent lectin inhibitors of the human immunodeficiency virus. [[PMID 25613831](#)]

A. Chatterjee et al. Griffithsin and Carrageenan Combination To Target Herpes Simplex Virus 2 and Human Papillomavirus. [[PMID 26369967](#)]

### Patent Status

- **U.S. Patent Filed:** U.S. Patent Application Number PCT/US2014/04099, Filed 05 Jun 2013

### Therapeutic Area

- Infectious Diseases