

Hello Bio, Inc.  
304 Wall St., Princeton, NJ 08540 USA

T. 609-683-7500  
F. 609-228-4994

customercare-usa@m2stage.hellobio.com



# DATASHEET

## JHU37152 (DREADD ligand)

### Product overview

<b>Name</b>	JHU37152 (DREADD ligand)
<b>Cat No</b>	HB6252
<b>Alternative names</b>	J52
<b>Purity</b>	>98%
<b>Description</b>	Novel DREADD agonist with high affinity and potency for hM3Dq and hM4Di. Active in vivo. Freebase.

### Images



### Biological Data

#### Biological description

#### Overview

JHU37152 is reported to be a novel DREADD agonist with high in vivo DREADD potency for CNS applications.

It has high affinity in vitro for hM3Dq and hM4Di ( $K_i$  values are 1.8 nM (hM3Dq) and 8.7 nM (hM4Di)).

It selectively displaces [ $^3$ H]clozapine from DREADDs and not from other clozapine-binding sites at concentrations up to 10 nM when tested for in situ [ $^3$ H]clozapine displacement in brain tissue from WT and  $D_1$ -DREADD mice.

JHU37152 activates hM3Dq and hM4Di with high potency and efficacy in fluorescent and BRET-based assays in HEK-293 cells ( $EC_{50}$  values are 5 and 0.5 nM at hM3Dq and hM4Di respectively).

#### Occupancy

JHU37152 exhibits high in vivo DREADD occupancy and was not reported to be a P-gp substrate.

#### In vivo application

JHU37152 is reported to be a potent in vivo DREADD agonist, which selectively inhibits locomotor activity in  $D_1$ -hM3Dq and  $D_1$ -hM4Di mice without any significant locomotor effects observed in wild type (WT) mice (at doses ranging 0.01 - 1 mg/kg).

It also produces robust and selective increases in hM3Dq-stimulated locomotion in rats expressing hM3Dq in tyrosine hydroxylase expressing neurons (at doses ranging 0.01 - 0.3 mg/kg).

While its selectivity is not ideal (i.e. comparable to clozapine), its high in vivo potency allows for dose adjustments with minimal off-target effects. The compound exhibits promising characteristics for

DREADD use in monkeys.

Water soluble version also available.

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## Solubility & Handling

### Storage instructions Solubility overview Important

Room temperature

Soluble in DMSO (100 mM)

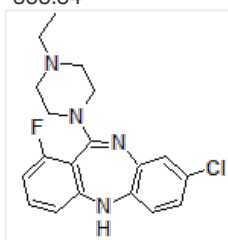
This product is for RESEARCH USE ONLY and is not intended for therapeutic or diagnostic use. Not for human or veterinary use

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## Chemical Data

### Chemical name Molecular Weight Chemical structure

8-chloro-11-(4-ethylpiperazin-1-yl)-1-fluoro-5H-dibenzo[b,e][1,4]diazepine  
358.84



### Molecular Formula CAS Number PubChem identifier SMILES Source InChi

C<sub>19</sub>H<sub>20</sub>ClFN<sub>4</sub>

2369979-67-7

0

CCN1CCN(CC1)C2=Nc4cc(Cl)ccc4Nc3cccc(F)c23

Synthetic

InChI=1S/C19H20ClFN4/c1-2-24-8-10-25(11-9-24)19-18-14(21)4-3-5-16(18)22-15-7-6-13(20)12-17(15)23-19/h3-7,12,22H,2,8-11H2,1H3

### InChiKey

NZMZJNNWMSYDNX-UHFFFAOYSA-N

### Appearance

Yellow solid

### Licensing details

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## References

### Chemogenetic ligands for translational neurotheranostics

Bonaventura et al (2018) bioRxiv doi: <https://doi.org/10.1101/487>

### High-potency ligands for DREADD imaging and activation in rodents and monkeys.

Bonaventura et al (2019) Nat Commun. 10(1)

#### PubMedID

[31604917](#)

### 0067 Humanized Chemogenetic Approach to Treat Sleep Apnea

Curado et al (2019) Sleep (42)

### OP-01-02 Graft-host synaptic connectivity can be chemogenetically inhibited with clinically relevant activators to eliminate graft-induced dyskinesias (GID) without losing anti-parkinsonian benefits of dopaminergic grafts

Subramanian et al (2019) World Congress On Parkinson's Disease And Related Disorders 2019 Poster abstract

### DREADDs: The Power of the Lock, the Weakness of the Key. Favoring the Pursuit of Specific Conditions Rather than Specific Ligands.

Goutaudier et al (2019) eNeuro 6

#### PubMedID

[31562177](#)

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