



RS 504393

Catalog No: tcsc0851

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## **Available Sizes**

Size: 5mg

Size: 10mg

Size: 25mg

Size: 50mg



## **Specifications**

CAS No:

300816-15-3

Formula:

 $C_{25}H_{27}N_3O_3$ 

**Pathway:** 

Immunology/Inflammation;GPCR/G Protein

**Target:** 

CCR;CCR

**Purity / Grade:** 

>98%

**Solubility:** 

DMSO : ≥ 10 mg/mL (23.95 mM)

**Observed Molecular Weight:** 

417.5

## **Product Description**

RS 504393 is a selective **CCR2** chemokine receptor antagonist ( $IC_{50}$  values are 89 nM and > 100  $\mu$ M for inhibition of human





recombinant CCR2 and CCR1 receptors respectively).

IC50 & Target: IC50: 89 nM (CCR2)<sup>[5]</sup>

In Vitro: RS 504393 inhibits the MCP-1-induced chemotaxis with an IC<sub>50</sub> of 330 nM. RS 504393 treatment suppresses allergen induced  $\beta$ -hexosaminidase release significantly. Without allergen priming, MCP-1 induces mast cell degranulation, which is completely suppressed by RS 504393<sup>[4]</sup>.

*In Vivo:* RS504393 (0.3-3  $\mu$ g) with CCL2 progressively blocks thermal hyperalgesia dose-dependently in mice<sup>[1]</sup>. RS 504393 (5 mg/kg, i.v.) supresses the elevated numbers of leukocytes and increased total protein content in BALF induced by The LPS. RS504393 significantly down regulates the LPS-induced elevation of IL-1 $\beta$ , PAI-1 mRNA and protein expressions. RS504393 significantly suppresses induced lung edema, protein-rich fluid, polymorphonuclear accumulation and bronchial wall thickening induced by LPS<sup>[2]</sup>. RS-504393 significantly reduces renal pathology, especially the extensive interstitial fibrosis mediated by decrease in type I collagen synthesis in a UUO model<sup>[3]</sup>.

All products are for RESEARCH USE ONLY. Not for diagnostic & therapeutic purposes!