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Effect of Pooled Human Intravenous Globulin (IVIG) on the Reversal of Cholinergic Inhibition of Smooth Muscle by Immunoglobulins (IgGs) from Patients with Scleroderma (SSc)

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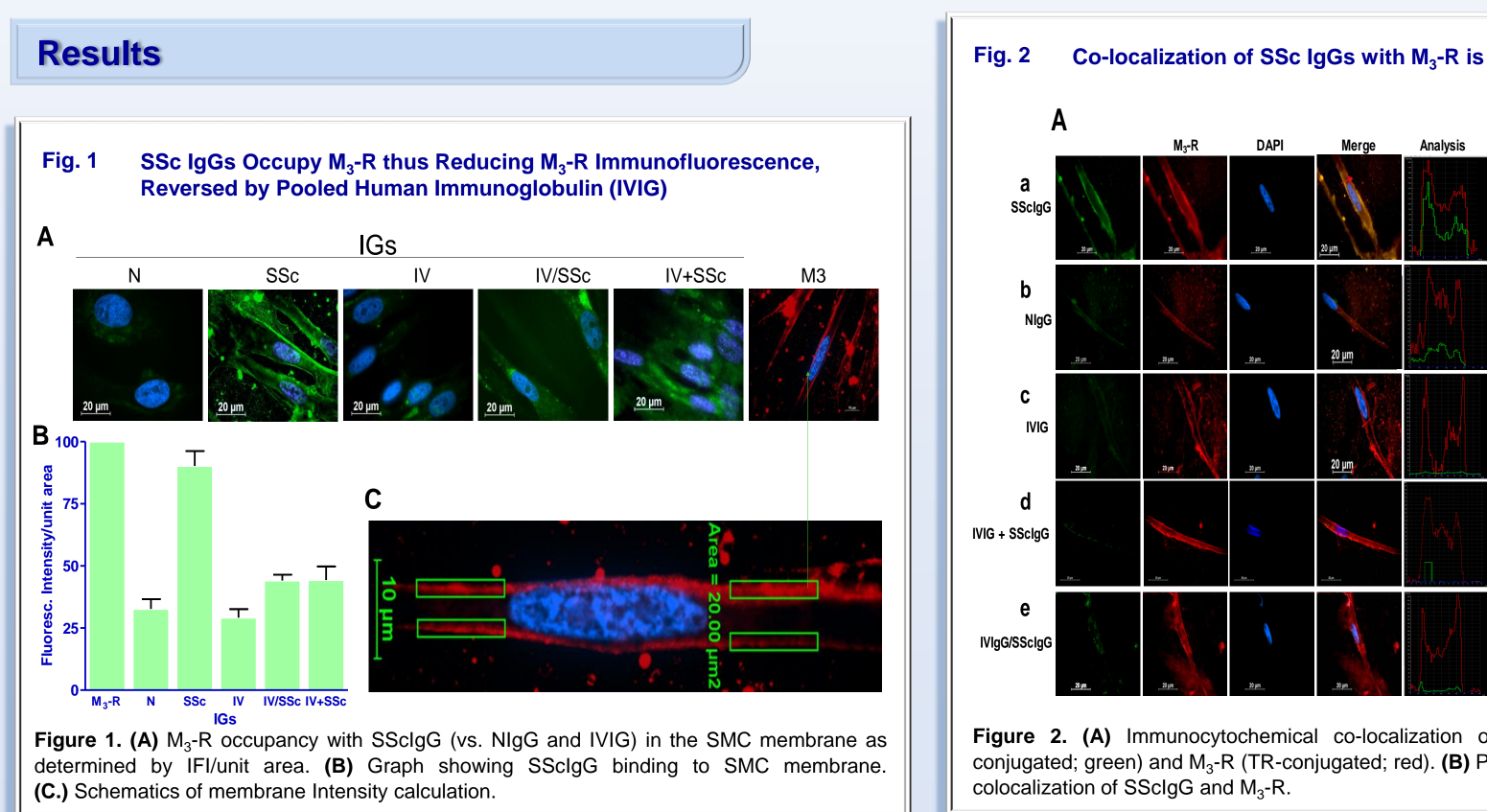
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Backgrounds & Aims

The gastrointestinal (GI) tract is the most common internal organ system affected in SSc. We and others have shown before that the SSc immunoglobulins (IgGs) cause selective blockade of muscarinic type-3 cholinergic (M₃-R) in the GI tract. Presently, there is no effective treatment for SSc although numerous cytotoxic and immunomodulatory agents have been employed with limited success and are marred with serious side effects. Present studies investigated the reversibility of SScIgGs-caused M₃-R blockade by the pooled Intravenous immunoglobulins (IVIG).

Methods

Effects of SScIgGs and IgGs from normal individuals (NIgGs) on M₃-R activation by bethanechol (BeCh) were determined in human internal anal sphincter (IAS) smooth muscle cells (SMCs), before and after IVIG. M_3 -R occupancy and binding by the SScIgG was determined via immunofluorescence (IF), ELISA, respectively. Functional Western blotting, and displacement of M_3 -R occupancy by the SScIgGs was determined employing different concentrations of the IgGs during the sustained phase of the BeCh-induced contraction of rat IAS smooth muscle strips.



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Beth 10⁻⁴ №

Time (min)

Figure 3. (A) SScIgG (and not NIgG and IVIG) in resemblance with darifenacin (M₃-R selective inhibitor) causes significant and conc.-dependent decrease in the BeCh-induced sustained contraction of IAS (*; p < 0.05; n = 6). (B) Actual tracings of the above effects.

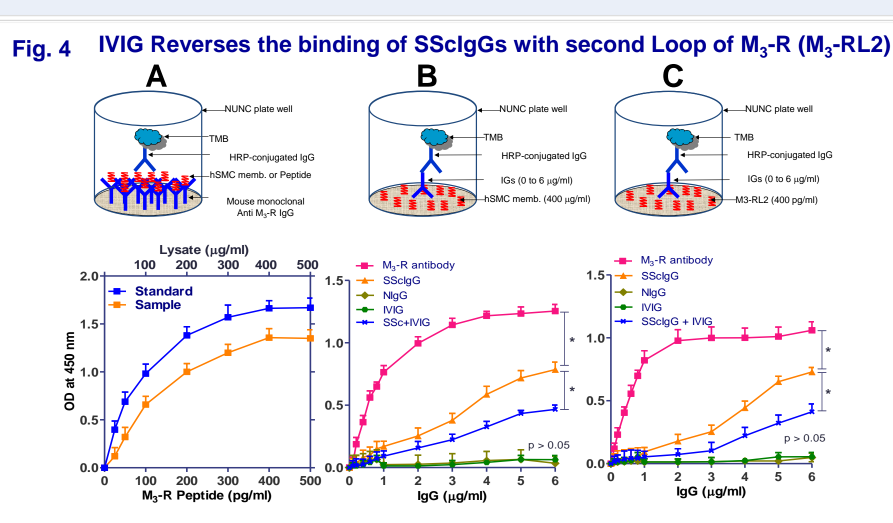


Figure 4. (A) ELISA binding studies for M₃-R peptide and human IAS SMC membrane fraction (HISMF): Standard curves with M₃-R and HISMF. **(B)** Data show M₃-R antibody and SScIgG bind with HISMF in a conc.-dependent manner (*; p < 0.05; n = 6), and IVIG significantly decreases this binding. (C) Similar data were obtained when M_3 -R peptide (M_3 -RL2) instead of HISMF was used.

Co-localization of SSc IgGs with M₃-R is Blocked by IVIG

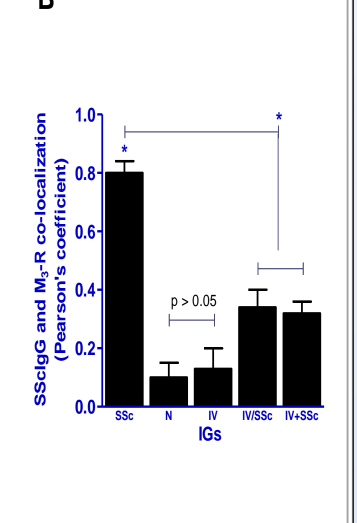
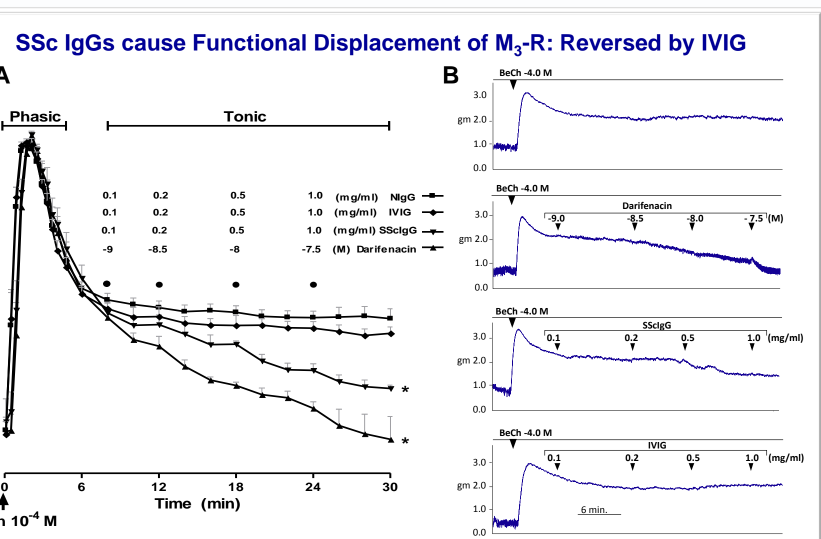
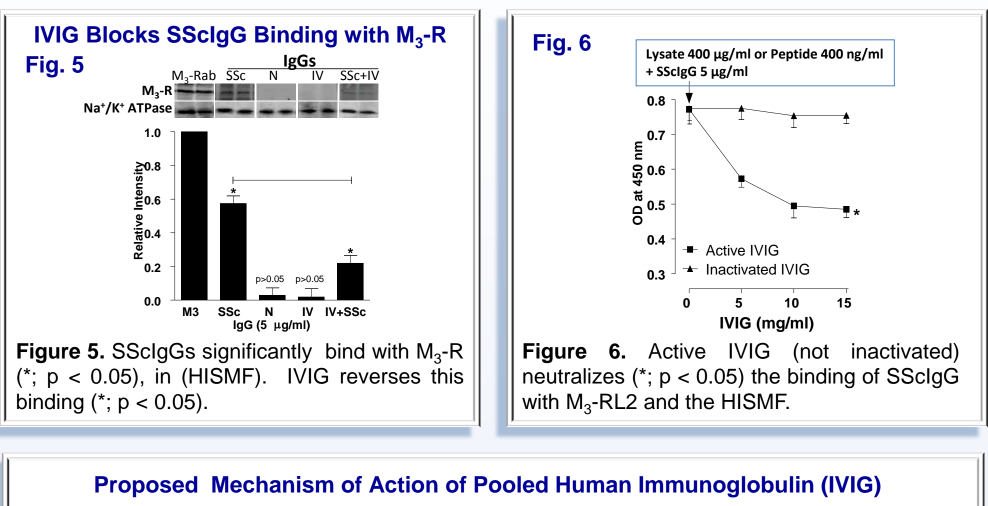
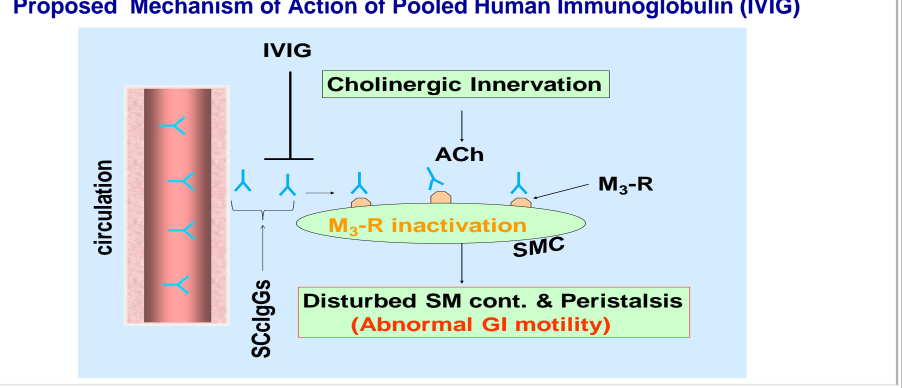


Figure 2. (A) Immunocytochemical co-localization of different IgGs (a,b,c,d,e) (FITCconjugated; green) and M₃-R (TR-conjugated; red). (B) Pearson's coefficient shows significant







Summary

- 1. IgGs from scleroderma patients (SScIgGs) inhibit muscarinic type-3 cholinergic (M₃-R) activation, as shown by the data in human IAS smooth muscle cells and rat smooth muscle strips.
- 2. SScIgGs inhibit M_3 -R occupation as shown by immunocytochemistry and Elisa-binding studies.
- 3. Pooled intravenous globulin (IVIG) reverses the M_3 -R occupancy and activation primarily by neutralizing circulating the SScIgGs.

Conclusions

- IVIG abrogates SScIgGs-mediated block of M₃-R by blocking the circulating SScIgGs.
- This mechanism may be partly responsible for the restoration of M₃-R-mediated cholinergic dysfunction in SSc-related GI manifestations.