

**Wykonawca:**

Aestimo s.c. [redacted]

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**Autorzy:**

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**Odpowiedź na pismo znak: MZ-PLA-460-16752-98/KK/14,**

zawierające uwagi Prezesa Agencji Oceny Technologii Medycznych do analiz załączonych dla produktów:

- **Rebif (interferonum beta-1a), roztwór do wstrzykiwań we wkładzie, 44 µg/0,5ml, kod EAN: 5909990728497;**
- **Rebif (interferonum beta-1a), roztwór do wstrzykiwań w ampułko-strzykawce, 44µg/0,5ml, kod EAN: 5909990874934,**

w ramach uzgodnionego z wnioskodawcą programu lekowego „Leczenie stwardnienia rozsianego (ICD-10 G 35)”.

W przedmiotowym piśmie Prezes Agencji Oceny Technologii Medycznych stwierdził, iż przedłożone analizy są niezgodne względem minimalnych wymagań określonych w rozporządzeniu Ministra Zdrowia z dnia 2 kwietnia 2012:

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**Analiza skuteczności**

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- Częstość rzutów choroby

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- Produkcja przeciwciał neutralizujących

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**Charakterystyka badania (*critical appraisal*)**

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**Podsumowanie**

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