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Thrombotic Microangiopathy Associated with Interferon Beta

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To the Editor: Interferon beta is a widely prescribed recombinant-protein therapy with a well-established favorable safety profile.¹ Here, we describe an unexpectedly high number of cases of thrombotic microangiopathy associated with severe or malignant hypertension in four patients with multiple sclerosis who were receiving therapy with recombinant interferon beta in South Scotland. A detailed review of the case histories of these patients, including a genetic analysis, did not identify any other causal factor for this condition, and an analysis of pharmacy records revealed a significant association with a common manufacturing source of interferon beta (Rebif, Merck) (Fig. 1, and the Supplementary Appendix, available with the full text of this letter at NEJM.org).

The regulatory authorities in the United Kingdom received six additional spontaneous reports of disorders related to thrombotic microangiopathy and interferon beta, all associated with the same manufacturer (for details, see the Supplementary Appendix). In December 2013, the Medicines and Healthcare Products Regulatory Agency (MHRA) issued a drug-safety update regarding a possible link between interferon beta and thrombotic microangiopathy.² In April 2009, the manufacturer of Rebif added a warning to the package insert about a possible association with thrombotic microangiopathy, which was specifically defined as the hemolytic–uremic syndrome and thrombotic thrombocytopenic purpura (HUS/TTP).

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We requested details regarding the geographical and temporal distribution of the HUS/TTP reports. Consistent with our observations in the United Kingdom, very few cases were reported globally in the first 9 years of safety monitoring. However, there has been a recent increase in reports from countries that share the same formulation as that used in the United Kingdom.³ Substantial caution is required in the interpretation of spontaneous reports of adverse drug reactions, since such reports are voluntary, are potentially subject to underreporting, and may be influenced by many factors, including the severity of the drug reaction, the diagnostic classification, and the degree of publicity. However, we did not detect such patterns or trends in the safe ty data from a similar recombinant interferon beta product, suggesting that further investigation of the association with changes in manufacturing may be worthwhile. We have raised these concerns with the manufacturer and regulatory authorities.

The patients in our analysis share important clinical features (for details, see the Supplementary Appendix). First, all the patients presented after years of well-tolerated treatment with inter-feron beta, making an association difficult to recognize. Second, fulminant presentation was associated with severe hypertension. Third, despite the emergency presentation, retrospective review identified chronic changes in all renal-biopsy specimens and a prodrome in three of the four patients with features that included newly diagnosed hypertension, hematologic abnormalities, and renal impairment in the months before diagnosis. We subsequently identified a similar prodrome in a recent fatal case in the United Kingdom. Thrombotic microangiopathy is therefore a serious and potentially fatal complication that has emerged late in the lifetime of recombinant inter-feron beta therapy. Early manifestations of this complication may be recognizable with increased vigilance, with the potential to mitigate severity.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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References

- PRISMS (Prevention of Relapses and Disability by Interferon beta-1a Subcutaneously in Multiple Sclerosis) Study Group. Randomised double-blind placebo-controlled study of interferon beta-1a in relapsing/remitting multiple sclerosis. Lancet. 1998; 352:1498–504. [PubMed: 9820297] Lancet. 1999; 353:678. Erratum.
- 2. Recombinant interferon-beta: thrombotic microangiopathy. [December 2013] Drug Safety Update. p. S3http://www.mhra.gov.uk/Safetyinformation/DrugSafetyUpdate/CON355481
- 3. Giovannoni G, Barbarash O, Casset-Semanaz F, et al. Safety and immunogenicity of a new formulation of interferon beta-1a (Rebif New Formulation) in a Phase IIIb study in patients with relapsing multiple sclerosis: 96-week results. Mult Scler. 2009; 15:219–28. [PubMed: 18755819]

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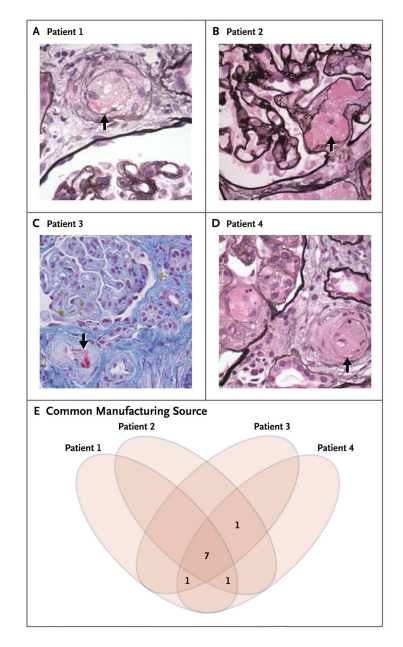


Figure 1. Cases of Thrombotic Microangiopathy Associated with a Common Manufacturing Source of Interferon Beta

Shown are renal-biopsy specimens obtained from four patients receiving recombinant interferon beta therapy in South Scotland in whom thrombotic micro-angiopathy was diagnosed during an 18-month period (Panels A through D). All four specimens show arteriolar luminal obliteration with swollen endothelium and fibrin (arrows; hematoxylin and eosin and silver staining [Panels A, B, and D] and Martius scarlet blue [MSB] staining [Panel C]). All four patients were treated with recombinant interferon beta from the same manufacturer, as confirmed by tracing of drug batches. The Venn diagram shows the overlap of 10 batches prescribed to patients in the year before the presentation of the index case (Panel E).

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