Ristocetin dependent cofactor activity in von Willebrand disease diagnosis: Limitations of relying on a single measure

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Abstract

Background: Von Willebrand disease (VWD) is a common inherited bleeding disorder, however the diagnosis can be complicated by a subjective bleeding history and issues with some current von Willebrand factor (VWF) laboratory assays.

Objectives: In the Zimmerman Program, we sought to determine how often a type 1 diagnosis was based on a single low VWF ristocetin cofactor (VWF:RCo) level result - coming from the common genetic variant p.D1472H or an isolated assay issue, if that low value was corroborated by the VWF glycoprotein-IbM (VWF:GPIbM) assay, and if retesting confirmed original levels.

Methods: New patients being evaluated for bleeding were consented. Analysis included VWF sequencing, bleeding scores, and comparisons of local VWF antigen (VWF:Ag) and VWF:RCo to central VWF:Ag and VWF:GPIbM.

Results: A total of 18% of VWD subjects had a low local VWF:RCo, but normal VWF:Ag and normal central testing including VWF:GPIbM. Seventy percent of the low VWF:RCo cohort had no pathogenic VWF variants; however, 33% carried p.D1472H. Low VWF:RCo subjects with follow-up local testing within 2 years showed those with p.D1472H continued to have low VWF:RCo and VWF:RCo/VWF:Ag ratio with normal VWF:GPIbM. Subjects without p.D1472H had an increase mean VWF:RCo, resulting in 59% with normal levels on repeat testing.

Conclusions: The diagnosis of VWD based on a single low VWF:RCo but normal VWF:Ag, was often attributed to p.D1472H or variability in VWF:RCo that was eliminated with VWF:GPIbM. Our study suggests that using VWF:RCo alone for diagnostic purposes may be insufficient while repeat VWF:RCo or VWF:GPIbM testing can be valuable in establishing a VWD diagnosis.

KEYWORDS
clinical laboratory tests, genetic polymorphism, Ristocetin cofactor, von Willebrand disease, von Willebrand factor
1 | INTRODUCTION

Von Willebrand disease (VWD) is a common inherited bleeding disorder, though diagnosis can be complicated because of a subjective bleeding history and issues with current von Willebrand factor (VWF) function assays. VWF activity is commonly measured in the United States using a ristocetin-dependent cofactor (VWF:RCo) assay, although this assay is highly variable and has poor sensitivity. The gain-of-function VWF glycoprotein-IbM (VWF:GPIbM) is a ristocetin-independent activity assay that provides both better precision and sensitivity and avoids the false low values seen with the common p.D1472H variant. p.D1472H causes decreased binding of ristocetin to the VWF A1 domain and has been previously identified in 63% African American and 17% Caucasian healthy controls; however, it is not associated with an increased bleeding risk.

In the Zimmerman Program on the Biology of VWD (Zimmerman Program) prospective study, we sought to determine how often a diagnosis of type 1 or low VWF was based on a single low VWF:RCo because of the p.D1472H variant or an isolated assay issue, if that low value was corroborated by the VWF:GPIbM assay, and if retesting confirmed original levels.

2 | STUDY DESIGN

New patients being evaluated for bleeding were consented and enrolled in the Zimmerman Program prospective cohort study at 12 hematology centers across the United States and Canada as previously described. The study was open to any patients who were new to the hematology clinic and being evaluated for a bleeding disorder. Informed consent was obtained from all subjects and the study protocol was approved by each local clinical acquisition center’s institutional review board. Hematology centers diagnosed subjects based on local laboratory values, and those with VWD continued in the study and were given the opportunity to participate in follow-up research blood draws. Central testing was performed at Versiti Blood Research Institute (BRI) including VWF antigen (VWF:Ag), VWF activity by VWF:GPIbM, and exonic VWF Sanger sequencing. Rare VWF genetic variants (present <1% of healthy controls) were considered potentially causative. The bleeding score was calculated according to the ISTH bleeding assessment tool (ISTH-BAT). Although the local centers had different normal ranges for VWF:Ag and VWF:RCo, a level of 50 was used as a cutoff in this analysis for type 1 VWD. GraphPad Prism was used to perform all statistical analyses. Comparisons of mean laboratory values and median bleeding scores used the nonparametric Mann–Whitney test.

3 | RESULTS AND DISCUSSION

A total of 1860 subjects were enrolled, including 677 with a new diagnosis of VWD and 1183 without VWD. A total of 454 VWD subjects with suspected type 1 or low VWF and complete laboratory and genetic results were analyzed (Figure 1). Most of this group was female (67%), self-identified as White (86%), non-Latino (86%), and was pediatric (69%) (Table 1). Eighty-two (18%) of these VWD subjects had a low local VWF:RCo (mean 43 IU/dl), but normal VWF:Ag (mean 62 IU/dl) and normal central testing with a mean VWF:Ag of 63 IU/dl (p = 0.9732) and mean VWF:GPIbM of 69 IU/dl (p < 0.0001) (Figure 2). This low VWF:RCo cohort had a median ISTH-BAT bleeding score of 3, which was not different from subjects without VWD (n = 1005; p = 0.2784), although there was a slight difference in age between these groups (median 12 vs. 14 years; p = 0.0195) (Table 2).

When we analyzed VWF sequencing results, we found that 70% of the single low VWF:RCo cohort had no potentially causative rare variant identified; however, 19 (33%) of these subjects did carry the common p.D1472H variant (17 heterozygous; 2 homozygous). Eight of these subjects had a reduced VWF:RCo/VWF:Ag ratio (<0.7) that could be considered type 2M VWD; however, all had normal VWF:GPIbM/VWF:Ag and no type 2M genetic variant identified. In the subjects with p.D1472H, 58% self-identified as White (including three Latino), 29% African American, and 10% Asian. Subjects with p.D1472H had a lower median bleeding score of 2 compared with 4 in those without p.D1472H (p = 0.2136) and those with a VWF variant (p = 0.2552); however, that difference was not significant.

In the Zimmerman Program prospective study, we hypothesized that repeat testing is critical for diagnosing VWD. To determine if subsequent retesting was valuable to confirm original findings, we examined a convenience sample including 37 of the 82 subjects (45%) with available follow-up testing performed locally and centrally within 2 years of the initial diagnostic draw (Figure 1). There were 15 subjects with p.D1472H who had follow-up testing where we observed a normal follow-up VWF:Ag in both the local and central laboratory testing. As expected, we found a low follow-up VWF:RCo (mean 461 IU/dl) similar to baseline, and a normal VWF:GPIbM (mean 65 IU/dl). Follow-up laboratory values were consistent with the presence of p.D1472H in
these subjects resulting in low VWF:RCO (Figure S1). The other 22 subjects without p.D1472H had a follow-up VWF:Ag (mean 671U/dl) that was similar to baseline and central testing. However, we observed an increase in mean VWF:RCO (571U/dl) compared with 441U/dl at baseline and a normal VWF:GPIbM consistent with baseline (Table 2). Retesting in this small sample group revealed an increase in mean VWF:RCO and a normal mean VWF:RCO/ VWF:Ag ratio (0.9), with approximately one-half of these subjects presenting with a normal VWF:RCO at follow-up.

Next, we identified 41 subjects who were enrolled in the study and categorized as non-VWD by their local hematology center, yet had a low VWF:RCO/VWF:Ag level <0.6 (Figure 1). All these
Subjects had normal VWF:Ag (mean 127 IU/dl), slightly reduced VWF:RCo (mean 63 IU/dl), and a normal VWF:GPIbM (90 IU/dl) and VWF:GPIbM/VWF:Ag ratio (0.9). Sequencing was performed where p.D1472H was detected in 71% (29) of these non-VWD subjects, which would explain their low VWF:RCo/VWF:Ag ratio. Twenty-one (72%) were heterozygous and eight (28%) were homozygous. Those with p.D1472H comprised 48% White, 38% African American, and 3% Asian participants. There was no difference in median bleeding score (4) between those with or without p.D1472H ($p = 0.6149$).

In addition to the small sample size, there are several limitations to this study. Although all new patients being evaluated for bleeding were approached for the study, we estimate that 75% agreed to participate, which could bias the composition of the cohort. Another limitation is the bleeding score may not accurately reflect all bleeding symptoms, especially in the pediatric population. It is also possible that genetic variants outside of VWF could be contributing to the phenotype in cases where the low VWF:RCo value could not be explained by p.D1472H or variability in the assay. Finally, in the non-VWD subjects with both a low VWF:RCo/VWF:Ag ratio and p.D1472H, clinical VWF exon 28
sequencing may have been performed and an erroneous VWD diagnosis avoided.

This report demonstrates that the reliance on VWF:RCo alone for diagnostic purposes can be insufficient, especially in those with the p.D1472H variant. Francis et al. showed similar results from a single institution study in which their retrospective analysis uncovered six patients diagnosed with type 1 VWD who had normal VWF:Ag levels and borderline low VWF:RCo, which could be explained by the presence of p.D1472H. Our data expand on this study and highlight the value of repeat testing along with using the VWF:GPIbM assay to avoid the potential pitfalls of the VWF:RCo assay. This study supports the recent VWD diagnostic guidelines which recommends using abnormal bleeding, VWF:Ag <50, and newer assays that measure VWF activity for diagnosing type 1 VWD. We also demonstrate that for patients who present with a disproportionate decrease in VWF:RCo compared with VWF:Ag, the presence of p.D1472H should be investigated to avoid misclassification as type 2M VWD.

In conclusion, 18% of our cohort had a VWD diagnosis based on a single low VWF:RCo but had normal VWF:GPIbM activity. The discrepancy between VWF:RCo and VWF:Ag was often because of the p.D1472H variant or variability in VWF:RCo that was improved with the VWF:GPIbM assay. Reliance on one VWF:RCo value alone for diagnostic purposes may be problematic, especially in those who have p.D1472H. Our findings suggest that performing VWF:GPIbM or repeat VWF:RCo testing can be valuable to ensure an accurate VWD diagnosis.

AUTHOR CONTRIBUTIONS

P. A. C. designed the research, analyzed data, and wrote the manuscript; U. O. S. performed research and analyzed data, S. L. H., V. H. F., and T. C. A. designed the research and edited the manuscript; R. R. M. conceived the original study, designed the research, and edited the manuscript. All authors have read and approved the final version of the manuscript.

ACKNOWLEDGMENTS

The authors thank all the subjects, hematology centers, and laboratory personnel who were all critical to this study. This research was supported by a grant from the National Institutes of Health NHLBI for the Zimmerman Program and multiple investigators under award numbers P01HL081588, P01HL144457, R01HL112614, R01HL136430, and R01HL126810. A complete list of Zimmerman Program investigators and contributing centers are included in the Appendix S1.

RELATIONSHIP DISCLOSURE

The authors do not have any relevant conflicts of interest to declare.

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SUPPORTING INFORMATION

Additional supporting information can be found online in the Supporting Information section at the end of this article.

How to cite this article: Christopherson PA, Haberichter SL, Flood VH, et al. Ristocetin dependent cofactor activity in von Willebrand disease diagnosis: Limitations of relying on a single measure. Res Pract Thromb Haemost. 2022;6:e12807. doi: 10.1002/rth2.12807