Clinical response to oral tofacitinib in pediatric patients with alopecia areata

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Key words: alopecia areata; JAK inhibitor; Janus kinase inhibitor; oral tofacitinib; pediatric population; treatment efficacy.

INTRODUCTION
Alopecia areata (AA) is an autoimmune disease characterized by cytotoxic T cell–driven inflammation targeting hair follicles and causing nonscarring hair loss. AA is associated with the overexpression of proinflammatory cytokines acting via Janus kinase (JAK)/signal transducers and activators of transcription signaling pathway. AA disproportionately affects youth and carries a significant psychosocial burden. While a myriad of treatments has been used for AA with varying success, there is no cure. Oral tofacitinib is a JAK3 inhibitor with established efficacy and tolerability in adult AA but not yet in pediatric AA. It is approved for juvenile idiopathic arthropathy (extended oligoarthritis, rheumatoid factor–positive or–negative polyarthritis, systemic juvenile idiopathic arthropathy without active systemic features) in patients as young as 2 years of age, with an acceptable safety profile. Baricitinib (JAK1 and JAK2 inhibitor) was recently Food and Drug Administration–approved for moderate-to-severe adult AA. We herein retrospectively report 10 chart-reviewed pediatric AA cases seen in the Columbia dermatology clinic and prescribed oral tofacitinib (Table I, Fig 1). As necessary precautions are required with JAK inhibitors (JAKis), baseline and follow-up laboratory testing (complete metabolic panel, glucose, complete blood count, tuberculosis screening, and lipid panel in those without a recent baseline) was performed. Patients were followed up for 2 years after hair regrowth to evaluate for relapse after dose tapering. Consent was obtained to use patient photographs (Fig 2).

CASES
Case 1
A 12-year-old boy presented for a 6-year history of patchy AA. He failed intralesional steroid (ILS) injections and topical clobetasol gel. On physical exam, the Severity of Alopecia Tool (SALT) score was 40%. He was prescribed 5 mg oral tofacitinib twice a day. After 1 month of treatment (electively taken once daily), he experienced full hair regrowth without side effects.

Case 2
A 13-year-old boy presented for a few month-history of alopecia universalis (AU). He had tried ILS, and with the understanding that this information may be publicly available.

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Funding sources: None.

IRB approval status: Reviewed and approved by Columbia University’s Human Research Protection Office IRB; Protocol # IRB-AAAU1635.

Consent for publication: Consent for the publication of patient photographs and medical information was provided by the authors at the time of article submission to the journal, and both the patient and parents gave consent for photographs and medical information to be published in print and online.

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JAAD Case Reports 2023;31:83-8.
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https://doi.org/10.1016/j.jdcr.2022.08.024
| Case | Sex | Age (yrs) | Medical history | AA subtype | Baseline SALT score (%) | Previous therapies | Tofacitinib Tx course | % change in the SALT score (% regrowth) |
|------|-----|-----------|----------------|------------|------------------|-------------------|---------------------|---------------------------------------|
| 1    | Boy | 12       | None           | Patchy     | 40               | Topical and ILS   | 1 5 (once daily)  | 0 100                                 | 1                                     |
| 2    | Boy | 13       | None           | AU         | 100              | Topical, oral, ILS| 1 10                | 0 100                                 | 1                                     |
| 3    | Girl| 13       | AD             | Patchy     | 60               | Topical and ILS   | 1 10                | 0 100                                 | 1                                     |
| 4    | Boy | 13       | None           | AU         | 100              | None              | 2 10                | 0 100                                 | 2                                     |
| 5    | Girl| 10       | None           | Patchy     | 50               | Oral and ILS      | 1 5 20              | 60 3                                  |                                       |
| 6    | Boy | 15       | Hashimoto's    | AU         | 90               | ILS, anthralin    | 2 10 80             | 11.1 4.5                              |                                       |
| 7    | Boy | 16       | None           | Patchy     | 50               | ILS               | 2.5 10              | 0 100                                 |                                       |
| 8    | Girl| 7        | None           | AT         | 100              | Tacrolimus ointment, ILS | 1 5 50 0 50 0 5                  |                                       |
| 9    | Girl| 12       | Celiac         | Ophiasis   | 90               | Topical and ILS   | 4 10                | 0 100                                 | 9                                     |
| 10   | Girl| 12       | None           | Patchy     | 95               | None              | 5 10                | 0 100                                 |                                        |

AA, Alopecia areata; AD, atopic dermatitis; AM, ante meridiem; AT, alopecia totalis; AU, alopecia universalis; ILS, intralesional steroids; PM, post meridiem; q, every; SALT, Severity of Alopecia Tool; Tx, treatment.
topical, and oral steroids without response. SALT score was 100%. He had positive scalp hair pull test, intact eyebrows and eyelashes, pitting nails, and patchy hair loss across bilateral arms and legs. He was therefore put on 10 mg oral tofacitinib BID. The patient presented after one month of treatment with full regrowth and no side effects other than mild facial acne.

### Case 3

A 13-year-old girl with atopic dermatitis presented for a 1-year history of rapid hair shedding unresponsive to topical steroids or ILS. SALT score was 60% and right eyelashes were substantially affected. After one month of well tolerated 10 mg oral tofacitinib BID, she experienced full hair regrowth.

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**Fig 1.** Patient timelines showing dates of initial presentation at our institution, therapies tried, oral tofacitinib initiation, and treatment response. †, Increased; AA, alopecia areata; AM, ante meridiem; Aug, August; Dec, December; ILS, intralesional steroids; Nov, November; q, every; Sept, September.
Case 4
A 13-year-old boy presented for a few months of functionally impairing AU. The SALT score was 80%. He had over 50% eyelash and eyebrow loss with severe eye irritation, no hair on arms, axillae, or legs, and significant nail pitting. He was therefore prescribed 10 mg oral tofacitinib twice a day. Two months later, he exhibited full regrowth without complaints.

Case 5
A 10-year-old girl presented for worsening patchy AA dating back to months prior to presentation. Prednisone and ILS were unsuccessful. Upon examination, the SALT score was 50% with eyebrow and eyelash involvement. After 1 month of 5 mg oral tofacitinib twice a day, she had regrown her eyebrows and coronal patches, achieving a 20% SALT score (ophiasis). The dose was hence doubled.
(10 mg twice a day). After 2 months, she experienced complete regrowth without adverse events.

**Case 6**

A 15-year-old boy with Hashimoto’s thyroiditis presented with a 1-year history of AU. He tried ILS and anthralin without amelioration. He reported severe eye irritation due to eyebrow and eyelash loss. On physical exam, the SALT score was 90%; alopecia barbae and patchy hair loss over bilateral arms were noted. After 2 months of 10 mg oral tofacitinib twice a day, the SALT score was down to 80% with some eyelash regrowth. After 2 and a half months of well-tolerated therapy, he presented with full hair regrowth.

**Case 7**

A 16-year-old boy presented for intermittent alopecia totalis (AT) that started before the age of 10 years. His condition was refractory to ILS and involved his eyebrows. The SALT score was 50%. He presented after 1 month of 5 mg oral tofacitinib twice a day with persistent scalp and left medical eyebrow patches of hair loss. After 2 months, he only regrew his eyebrows and eyelashes. The patient’s tofacitinib dose was thus increased to 10 mg twice a day. The following month, he started to regrow fine scalp hair. One month later, he presented with full regrowth and no complaints.

**Case 8**

A 7-year-old girl presented for AA that started a few months ago (Fig 2, A) along her occipital hairline with gradual worsening to AT. She was applying tacrolimus ointment without improvement. She couldn’t tolerate ILS given the large area of involvement. She reported eyelash loss that had mostly regrown. The mother was unsure if the patient grew body hair prior to disease onset. SALT score was 100%. The patient was initially put on 5 mg oral tofacitinib BID for one month with minor regrowth. She was henceforth prescribed 5 mg in the morning (AM), 10 mg in the evening (PM) for a month. She presented with a SALT score of 70% and normal eyebrows and eyelashes. She was subsequently put on 10 mg BID and returned after four months with full regrowth (Fig 2, B) and no side effects.

**Case 9**

A 12-year-old girl with celiac disease presented for deteriorating AA that first appeared at the age of 4 years. She reported losing patches of eyebrows and eyelashes with severe chronic ocular inflammation. She tried clobetasol and ILS without response. On examination, the SALT score was 90% (ophiasis pattern) and arms and legs were hairless. After 3 months on 10 mg oral tofacitinib twice a day, the SALT score was reduced to 20%. She achieved full regrowth after 6 months without side effects.

**Case 10**

A 12-year-old treatment-naive girl presented for a 1-year history of severe patchy AA. The SALT score was 95%. After 4 months on 5 mg oral tofacitinib twice a day, the SALT score was reduced to 10% (ophiasis). After 5 months on 10 mg oral tofacitinib twice a day, she achieved full regrowth and was doing well.

**DISCUSSION**

Oral tofacitinib has been gaining momentum among JAKis for the treatment of adult AA. Studies investigating its use for pediatric AA are scarce. Craiglow et al reported positive outcomes with 5 mg oral tofacitinib twice a day for pediatric AA.

Our patient population was equally distributed with respect to sex but substantially varied in terms of AA onset (as early as the age of 4 years, latest onset at the age of 14 years), disease subtype (5/10 patchy AA, 1/10 AT, 1/10 ophiasis, 3/10 AU) and severity (SALT scores ranging from 40% to 100%), and eyebrow (6/10) and eyelash involvement (7/10). Moreover, 3 of 10 had a concomitant autoimmune disease (eczema, Hashimoto’s thyroiditis, celiac) known to be associated with AA. There was no family history of AA or other autoimmune diseases. Prior attempted treatments included topical steroids (4/10), oral steroids (2/10), ILS (8/10), anthralin (1/10), and tacrolimus ointment (1/10). Two patients were treatment naive. One patient responded to a dose of 5 mg once daily. Four patients had their dose tapered up from 5 to 10 mg twice a day due to incomplete response, 1 of whom was bridged with 5 mg in the AM and 10 mg in the PM prior to the 10 mg twice a day dose step-up. Remaining 5 patients were on 10 mg twice a day from the start. Tofacitinib was tapered off for all after achieving full hair regrowth without any relapse.

Jerjen et al proved tolerability and clinical improvement with oral tofacitinib in preadolescents with AA. Likewise, all of our 10 patients achieved full response with variable treatment duration. Our findings are also supported by a recent pediatric case series where 8 of 11 patients experienced hair regrowth on oral tofacitinib (5-10 mg twice a day) without any adverse events. Similarly, none of our patients developed adverse events or laboratory abnormalities on follow-up testing. One case developed mild facial acne which subsided at the end of the treatment course. Patients with depression and
social anxiety reported improvement after tofacitinib therapy.

A case series found topical tofacitinib to be a reasonable adjunct (second-line) treatment for pediatric AA in whom systemic therapies are contraindicated. This was supported by a case report of localized AA (eyelashes), whose authors concluded that while topical JAKis spare patients the side effects of systemic therapies, they are impractical and unlikely to achieve adequate hair regrowth in extensive AA.

Our case series emphasizes the safety and efficacy of off-label oral tofacitinib in pediatric AA patients, regardless of age, presence of other autoimmune diseases, disease duration, subtype, or severity. The higher dose (10 mg twice a day) could explain the quick and full response achieved by all patients. Limitations include small sample size, retrospective nature, lack of the control group, and short follow-up period, which precludes long-term data extrapolation. As such, larger, randomized, controlled trials comparing tofacitinib to other JAKis (eg, baricitinib) are needed to fill the knowledge gap in pediatric literature. Our results are promising and pave the way for the determination of the safe and effective dose of oral tofacitinib for long-term pediatric AA management.

Conflicts of interest
None disclosed.

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