A 37 year-old male, who identified as a 'body builder,' was admitted under the acute medical assessment unit with complaints of sudden-onset shortness of breath, which was constant in nature and associated with 'tightness' in the throat. On examination he was afebrile but had laboured breathing. He was also noted to have a hoarse voice. These symptoms were sudden onset in nature and had been present for several hours prior to presentation. Oxygen saturation was 100% on room air while respiratory and heart rate were 25/min and 76/min, respectively. Neurological and abdominal examinations were unremarkable. Complete blood count, arterial blood gases, electrocardiogram, and chest x-ray were all deemed to be normal. History revealed that the patient regularly took the anabolic steroid Deca Durabolin® (nandrolone decanoate) and had done so for four years prior to admission. The patient admitted to injecting AAS once per week. The patient denied taking any other medications. Interestingly, the patient reported that he had experienced similar symptoms two days earlier and attended his primary care physician for advice. He was prescribed a salbutamol inhaler but reported that this offered minimal relief of his symptoms. Additionally, he had experienced similar symptoms intermittently whilst working out at the gym. A provisional diagnosis of an allergic reaction was made. The patient improved with administration of chlorphenamine and was discharged.

Ten days later, he was admitted to the ear, nose, throat (ENT) emergency clinic with worsening of ongoing symptoms, which were characterised by deteriorating breathlessness on minimal exertion. Subjectively, breathy and hoarse voices were noted. More concerning, marked stridor was also present. Salbutamol, antihistamines, prednisolone and amoxicillin offered minimal relief. Nebulised adrenaline (1ml 1:1000) was prescribed for stridor and further examination was performed by means of nasal endoscopy which showed no superior mediastinal mass or neck mass. Subsequently, a computerised tomography scan of his neck and chest was performed which showed no superior mediastinal mass or neck soft tissue mass. No other cause of vocal cord palsy was identified. He significantly improved the next morning and upon discharge, his voice was almost back to normal. He denied any other ongoing throat symptoms. He was strongly advised against the use of anabolic steroids and was informed about his underlying problem.

Three months later, he was seen in the otolaryngology outpatient clinic where he confirmed avoiding taking AAS. His breathing had significantly improved and his voice was almost back to normal. He denied any other ongoing throat symptoms. He was not keen for reassessment of his vocal cords by flexible laryngoscopy. He was discharged from the clinic.
Discussion

Steroid abuse, that is consuming supraphysiological amounts of anabolic androgenic steroids (AAS) for non-medical purposes, poses a significant public health problem with more than three million Americans currently misusing AAS. Anabolic steroids remain the most common pharmacological agent detected by antidoping tests in the summer Olympic Games [19]. Indeed, the last 10 years have seen an increased prevalence of AAS, particularly among adolescents and athletes [3]. However, it is thought that most AAS abusers today are non-elite sportsmen (recreational gym attendees) using the drugs for cosmetic purposes. Evidence is now emerging regarding the increasing problem of muscle dysmorphia or so called 'bigorexia' in males [20]. Muscle dysmorphia is defined as a pre-occupation with increasing muscles mass or leanness. This body-related anxiety often results in men turning to AAS to achieve their perceived goals.

In the United Kingdom, AAS are controlled as Class C substances under the Misuse of Drugs Act 1971. Consequently, AAS are often acquired from unreliable sources and the precise steroid agent and associated side effects are often unknown prior to consumption. It has been reported that AAS misusers can consume 10 to 100 times the recommended therapeutic dose of steroid leading to significant adverse effects [21]. Reported side effects of AAS misuse include increased sexual drive, development of acne, increase in body hair, and heightened aggression. Additionally, testicular atrophy and a decrease in sperm quality and quantity may occur secondary to suppression of gonadotropin production. Furthermore, AAS are known to be a risk factor for cardiovascular complications including myocardial infarction, development of cardiomyopathy, and sudden cardiac death [14].

Anabolic steroids for medical use, a change in voice can be seen in up to 10% [22]. A 'safe dose' has not yet been reported in order to avoid such symptoms [23]. The most noticeable change is a lowering in frequency of the voice as well a marked lowering in pitch [24]. Talaat et al. [22] examined histological changes in the larynx of female mice that had been treated with AAS for up to three months. Reversible changes included vascular congestion, oedema and perivascular and periductal cellular infiltration [22]. Irreversible changes included squamous metaplasia and parakeratosis of the epithelium and hypertrophy of the muscle fibers of the thyroarytenoid muscle [22]. At the time of this writing, no reports could be found of structural or histological laryngeal changes in males.

In our case, we present a sudden onset vocal cord palsy in a 37-year-old man who admitted to chronic anabolic steroid misuse. Vocal cord palsy is a relatively common clinical occurrence seen in 0.42% of the population [25]. The vocal cords are innervated by the recurrent laryngeal nerves, which can be damaged secondary to local malignancy, surgery, intubation, or idiopathic causes [26]. Gerritsma et al. [23] examined virilisation of the voice in post-menopausal woman taking AAS and suggested that changes in the rigidity of the vocal folds secondary to an increase in muscle deposition may be the cause of the lower vocal frequency commonly seen [23]. One could hypothesize that increased rigidity could ultimately lead to vocal cord paralysis given the extremely high amounts of AAS taken. Alternatively, vocal cord paralysis could occur due to damage to the recurrent laryngeal nerves. Studies have demonstrated AAS to be neurotoxic when injected in the proximity of peripheral nerves [27]. Although the exact mechanism by which this occurs is poorly understood, it is postulated that AAS initiate the apoptotic cascade of neuronal cells [28].
Vocal Cord Paralysis Secondary to Anabolic Steroid Abuse


Conclusion

AAS misuse is common and associated with a plethora of adverse effects. Due to the illicit nature of AAS, evidence of such side effects often go undocumented and are limited to case reports. Given the scale of AAS it is important that interesting complications of these pharmacological agents are reported. This case report serves to demonstrate the need for clinicians to consider AAS in the differential of a young, athletic male presenting to the ER or ENT clinic without a clear etiology for vocal cord palsy.

References
