Some rewarding effects of androgens may be mediated by actions of its 5α-reduced metabolite 3α-androstanediol

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Abstract

The abuse of anabolic–androgenic steroids (AS) is a growing problem; however, the effects and mechanisms underlying their addictive effects are not well understood. Research findings regarding androgen abuse in people and hedonic effects of androgens in laboratory rats are reviewed. Androgens, like other steroids, can have traditional actions via cognate intracellular steroid receptors, as well as other substrates. Our recent results indicate that testosterone (T) metabolites may have actions in part via γ-aminobutyric acid (GABA)A/benzodiazepine receptor complexes (GBRs) and/or dopaminergic neurons in the nucleus accumbens, to mediate T’s positive hedonic states. This may provide the basis for positive reinforcing effects of androgen seeking and use behavior. Following a comprehensive review of the background literature, our findings are presented that have explored the extent to which metabolites of T mediate euphorogenic effects of androgens by acting in the nucleus accumbens. Then results regarding whether GBRs are necessary substrates for androgens’ positive hedonic effects are discussed. Lastly, research that addresses if dopaminergic neurons in the nucleus accumbens are necessary for these effects of androgens are discussed. This review provides a comprehensive examination of the hedonic properties and abuse/addiction potential of androgens and the putative mechanisms underlying these effects.

Keywords: Testosterone; Reward; Reinforcement; Hedonic; Conditioning; GABA_A receptors; Dopamine receptors; Anxiety; Affect; Learning

1. Introduction—overview

Anabolic–androgenic steroids (AS) are the synthetic variants of the primary masculinizing androgen, testosterone (T). They are abused by growing numbers of individuals in this country ranging from adolescents, seeking to improve their appearance, to professional athletes attempting to elevate their performance. The costs associated with AS abuse are substantial. For the individual, AS abuse is associated with many adverse physical and behavioral consequences. For society at large, AS abuse has spawned a significant black market which has promoted criminal behavior and placed strain on law enforcement agencies. The aim of this review is to explore more fully the causes of androgen abuse in people by describing research from our laboratory and others investigating hedonic effects of androgens.

In contrast to our understanding of the classic drugs of abuse like cocaine, heroin and alcohol, relatively little is known about the causes of AS abuse. Indeed, the principal question surrounding most drugs of abuse is to what extent they produce euphorogenic (positive hedonic and rewarding) effects, which can maintain and/or exacerbate future drug seeking behavior. Our earlier findings, and those from other laboratories, are mixed. While positive hedonic effects of T have been reported by some, the results have been compromised by serious procedural problems and significant variability. Our recent research suggests that a portion of this variability may be due to differences in the metabolism of T and the resulting availability of androgen metabolites at receptor sites in the brain. Our results suggest that the actions of T metabolites, perhaps in part through γ-aminobutyric acid (GABA)A/benzodiazepine receptor complexes (GBRs) and/or dopaminergic neurons in the nucleus accumbens (NA), may mediate T’s production of positive hedonic states and consequently provide the basis for positively reinforcing effects on drug seeking and use behaviors.

This review paper initially summarizes background research that has prompted our investigations aimed at answering four key questions geared towards elucidating the mechanism of androgens’ actions in the NA to produce hedonic effects. First, we...
have investigated whether the ability of T to produce positive hedonic effects involves its metabolites. Comparisons between T and its metabolites and effects of pharmacological blockade of T metabolism have been used to ascertain whether such manipulations inhibit the positive hedonic effects produced by T. Second, because some of T’s metabolites do not bind well to intracellular androgen receptors (ARs), we have examined effects of blocking ARs, on hedonic effects of androgens. Third, the importance of GBRs in the NA as substrates for androgens’ hedonic effects have been addressed using intra-accumbens infusions of antagonists and agonists to investigate the importance of GBRs for androgens’ production of positive hedonic state. Fourth, whether dopaminergic neurons in the NA are required for androgens’ hedonic effects has been investigated using intra-accumbens 6-OHDA infusions. These questions have been examined in rats by using classic methods of behavioral pharmacology and neuroendocrinology to manipulate androgens’ actions and examining hedonic effects using a traditional Pavlovian conditioning technique, conditioned place preference (CPP), to assess the ability of androgens to produce positive hedonic states. The results of this research from our laboratory comprise the second half of this review paper. Taken together, the information presented in this review expands our understanding of the biochemical pathways through which androgens work to produce euphorogenic effects which may lead to their abuse.

2. Background and significance definition of AS

Anabolic steroids are androgenic and/or synthetic variants of T, which is the primary masculinizing hormone secreted by the mammalian testes, and is also secreted by ovaries and adrenals, albeit in lower concentrations. The original goal in producing these drugs was to promote the well-known anabolic (tissue building) effects of T without its androgenic (masculinizing) properties. Importantly, although the androgenic effects of some of the presently marketed anabolic steroids have been reduced, they have not been eliminated. Therefore, to date, no exclusively anabolic substance has yet been found (Bahrke et al., 1996).

2.1. Medical use of AS

It has been asserted that the German government under Hitler developed and were the first to use AS, allegedly in an attempt to create an army of supermen (e.g., Marshall, 1988). After the war, AS were used in legitimate medicine to treat female breast cancer (by reducing estrogen), to combat two different forms of anemia, and to reduce the effects of hereditary angioedema (Wright, 1980; Bahrke et al., 1998). While other drugs currently are more effective in initiating delayed puberty in preadolescents, growth promotion, treatment of micropenis, and treatment of hypogonadism (Moore, 1988; Wilson and Griffin, 1980; Bahrke et al., 1996).

2.2. Elicit use of AS

As the medical need for AS declined, their popularity among athletes soared dramatically. Soviet weight lifters began using them as power boosters in Olympic competitions in the 1950s, followed by the Americans in the 1960s (Wade, 1972; Yesalis and Cowart, 1998). Recent confirmations indicate the East German government sanctioned a massive experimental program in which their top athletes were administered AS for years in order to enhance athletic performance and medal counts in the Olympic games (Franke and Berendonk, 1997).

2.3. Incidence of AS use among athletes

While AS initially were used by super-athletes as a means for adding an edge to a performance already close to perfection, the word gradually spread that they could be effective for any sport that required strength. Although actual figures on the incidence and prevalence of AS use among elite, amateur, and recreational athletes is just beginning to emerge, it is generally agreed that AS currently are used widely by professional, college, and high school football and baseball players, shot-putters, discus throwers, swimmers, sprinters, tennis players, and bicyclists (Yesalis and Cowart, 1998; Bahrke et al., 1996). For example, studies commissioned by the National Collegiate Athletic Association (NCAA) (Anderson et al., 1991, 1999), indicate that roughly 5% of male and female athletes reported using AS, with the highest user rates (10%) seen in collegiate football players. In a survey of track and field athletes in the 1972 Olympics, 68% reported prior steroid use (Silvester, 1973). Among elite power lifters, 55% of those interviewed conceded prior AS use (Yesalis et al., 1988). In a study of amateur competitive body builders, over half of the men and 10% of the woman reported that they had used AS at some point in their life (Tricker et al., 1989).

2.4. Effects of AS use among athletes

Although AS are widely used today to enhance strength and athletic performance, there is little scientific documentation of their reputed benefits. An older review concluded that AS were beneficial only to enhance strength and athletic performance if the individuals were in a continuing program of intensive exercise coupled with a high protein diet (Haupt and Rovere, 1984). However, a more recent study (Bhasin et al., 1996) controlled dietary factors, type of exercise, weight lifting experience, and duration and dose of AS exposure and was the first to demonstrate that supraphysiological dosages of T, with or without strength training, increase fat-free mass, muscle size, and strength in normal men. The results further showed that men who received 10 weeks of strength training or 10 weeks of T experienced significant increases in muscle size and strength; however, those that received both T and strength training had increased strength, tricep and quadriceps size and fat-free mass that was well beyond that of men who received placebos, just AS with no strength training, or just strength training with no AS. These data demonstrate that T and exercise produced additive increases in performance. Also, a review of confiscated documents following the collapse of the German Democratic Republic clearly shows that AS had a positive effect on athletic performance in adult women and children (Franke and Berendonk, 1997).
A credibility and information gap between athletes and the medical/scientific community has resulted in the banning of AS by a number of athletic organizations (cf., Haupt and Rovere, 1984; Yesalis and Cowart, 1998) so as to protect athletes from potentially harmful side effects (see below). While sophisticated detection procedures have been developed in order to enforce these bans, athletes continue to abuse AS, often times by utilizing masking agents, or new AS which as yet cannot be detected.

2.5. Incidence of AS use among adolescents

Anabolic–androgenic steroids abuse is no longer confined to elite Olympic, professional, college or high school athletes. Lifetime use incidence is 5% for males and 3% for females, which indicates that more than 1 million Americans have taken AS illicitly. Although it may not be surprising that 55% of 27-year-old male and 10% of 24-year-old female body builders admit to using AS, and that the incidence in college athletes is estimated at 20%, AS abuse is now a problem that influences a much broader population including adolescents and young adults. The first nationwide study of AS abuse in 1987 particularly is revealing (Buckley et al., 1988) since other more recent studies generally confirm these findings. The study showed that 7% of 12th grade male and 3% of female students were using or had used AS and that two thirds of the user group initiated use when they were 16 years of age or younger. Importantly, the survey also showed a number of intriguing trends in AS abuse. 47% of users reported that the main reason for using the drug was to improve athletic performance but an alarming 27% of the user group listed appearance as the main reason. Also, many users in the study could be described as habitual because (1) 40% of the self-identified users reported using AS for five or more cycles with each cycle usually lasting six to 12 weeks, and (2) 44% of users responded that they used more than one AS drug at the same time (“stacking”). More recent surveys of 9th through 12th grade public and private high school students confirm the earlier results. For example, the 1995 Youth Risk and Behavior Surveillance System data showed that 4.9% of boys and 2.4% of girls have used AS at least once in their lives (Kann et al., 1996). More recently, it was reported that 4% of Massachusetts junior high school students have used AS (Faigenbaum et al., 1998). These findings indicate that the epidemiology of AS abuse is following a classic pattern similar to that of other drugs of abuse. Namely, abuse by elite groups is soon followed by widespread use of the drug by many other segments of the general population. Extrapolating from all of these findings, it is estimated that approximately 375,000 adolescent boys and 175,000 adolescent girls are steroid users. This represents a significant population at risk as the potential for adverse effects (see below) may be greatest in this age group.

2.6. Traffic in AS

The alarming figures reported above are even more striking in view of legislation in 1990 (Anabolic Steroids Control Act) that attempted to control by reclassifying AS in the United States as Schedule III controlled substances. Distribution without a prescription is a felony punishable by up to 5 years in prison and a $250,000 fine. Possession of AS in the US is a misdemeanor punishable by up to 1 year imprisonment and a minimum of a $1000 fine. AS use is a pervasive international problem (other countries are reporting problems similar to those seen in the U.S. (Bahrke, 1995).

Despite government attempts to control AS, their demand has resulted in rapidly escalating black market, which was estimated to involve $500 million in sales in 1994 (Kouri et al., 1994). A growing concern is the nutritional supplement field in which products such as androstenedione (Andro) and dehydroepiandrosterone (DHEA) contain AS. Not covered by the Anabolic Steroids Control Act, over-the-counter and internet sales of Andro and DHEA represent a sizable and growing portion of the $15 billion nutritional supplement market (Yesalis and Cowart, 1998).

2.7. Health risks associated with AS

The health risks associated with AS abuse are considerable (e.g., Haupt and Rovere, 1984; Yesalis and Cowart, 1998; Bahrke et al., 1996; Yesalis and Bahrke, 1995). The androgenic (masculinizing) risk is particularly acute for women, in whom it may be impossible to reverse masculine traits, such as facial hair, deepening of voice, and male physique, once they appear. In adult males, AS abuse can lead to kidney and liver damage, liver cancer, heart disease, and hypertension. It also can cause suppression of T production, enlargement of mammary tissue (gynecomastia), testicular atrophy, and decreased spermatogenesis. In adolescent males, AS abuse can hasten the onset of adulthood, promote early baldness, limit stature, and cause premature growth plate closing.

Many studies indicate that androgens modulate aggressive and copulatory behavior in rodents (Svare et al., 1983; Kinsley and Svare, 1988; Frye et al., 1996a,b). There are far fewer studies documenting a positive relationship between endogenous androgens and aggressive and copulatory behavior in humans. However, in men, there is ample evidence for an association between androgens and these behaviors (Bahrke et al., 1990, 1996).

2.8. Hedonic effects of androgens among people

There is now little doubt that AS can have significant effects upon mood and mental disorders (cf., Bahrke et al., 1990, 1996). Clinical reports showing increases in affective and psychotic syndromes, a number of which are very violent and suicidal in nature, are associated with AS use in individuals seeking to improve their performance or appearance (e.g., Pope and Katz, 1987, 1988; Pope et al., 1994). Indeed, there are now several legal cases in which defendants have claimed that AS’ effects upon their behavior promoted their criminal acts (e.g., Bahrke et al., 1990). Data from the National Household Survey on Drug Abuse have shown a strong association between AS use and self-acknowledged acts of violence against people and property crimes. Based upon this accumulating information, some researchers (Orchard and Best, 1994) have suggested that
violent offenders should be tested routinely for AS so as to further document the relationship and develop methods to control steroid abuse.

Case studies and anecdotal reports show AS abuse has been associated with changes in depression, euphoria, hypomania, increased aggression, libido, alertness, irritability, anger, anxiety, energy, hostility, mood swings, psychotic episodes, and violent rages (cf., Bahrke et al., 1998; Yesalis and Cowart, 1998). Self-reported changes in mood, behavior, and somatic perceptions also have been associated with AS abuse (Bahrke et al., 1992; Wilson, 1988). Up to 43% of AS users report feeling "high" or feeling extreme pleasure from using AS over extended periods of time (Brower et al., 1991). Similarly, many other uncontrolled studies have reported euphoric effects among athletes who have taken AS (reviewed in Taylor, 1987). Interestingly, some studies show that AS elicit electroencephalographic changes similar to those seen with amphetamines and tricyclic antidepressants (Bahrke et al., 1990). Indeed, these findings are consistent with reports that T was used to treat depression in the 1930's (Altschule and Tilletson, 1948).

2.9. Dependence

Some researchers have raised the possibility that dependence may result from prolonged AS abuse (Tennant et al., 1988; Wright, 1980; Kashkin and Kleber, 1989). They note that AS abusers often experience a stimulant-like withdrawal syndrome characterized by depressive symptoms. Moreover, a number of studies and case reports have documented behavior, perceptions, and attitudes in some AS abusers that are indicative of dependence (e.g., Brower et al., 1990, 1991, 1989; Corcoran and Longo, 1992; Pope and Katz, 1994). Since 1988, there have been at least four case reports of AS dependence in the medical literature. There is the case of a 24-year-old noncompetitive weight lifter, who met 6 of the 9 DSM criteria for dependence and experienced suicidal depression upon cessation of AS use (Brower et al., 1989). Another 22-year-old noncompetitive weight lifter reported low self-esteem and AS cravings so severe after AS cessation (Hays et al., 1990) that he was unable to discontinue their use. As well, 3 weightlifters report initiating AS use to enhance performance but maintaining use to prevent withdrawal (Tennant et al., 1988). A 30-year-old woman was also described who had been taking AS for 4 years; she met 5 of 7 of the DSM criteria for dependence (Copeland et al., 1998).

These findings suggest that the positive hedonic effects of AS are very powerful and may be a primary mitigating factor for their continued use. Findings from survey research are consistent with the notion of dependence as demonstrated by intention for continued AS use despite adverse consequences (a DSM criteria for dependence). One-fourth of high school seniors that admitted to using AS indicated that they would not stop using them even if the drugs led to permanent sterility, increase in the risk of cancer, or heart attacks (Yesalis et al., 1990); the response rate for heavy users of AS was as high as 50%. The incidence of AS users that meet the DSM criteria for abuse or dependence are as high as 100% and 75%, respectively (Brower et al., 1991, 1990). Other studies that have used the DSM criteria for substance dependence report rates of 15 to 69% (Malone et al., 1995; Pope and Katz, 1994; Clancy and Yates, 1992).

2.10. Hedonic effects of androgens in animal models

While these findings are interesting, their unsystematic and anecdotal nature limits their significance in understanding the mechanisms underlying AS dependence. Evidence from animal studies suggests T can have positive hedonic effects. Testosterone (Olds, 1958; Caggiula, 1970; Campbell, 1970), like many drugs of abuse (Kornetsky, 1995), will increase rates of bar pressing for electrical brain stimulation, which is considered an indication of a drug's rewarding effects. AS administration will increase the rate of bar pressing to deliver electrical brain stimulation to the mesolimbic system (Kornetsky, 1995; Caggiula and Hoebel, 1966; Herberg, 1963; Clark et al., 1996). As well, animals can be made to be physically dependent on AS (Bonson et al., 1994). Male hamsters preferentially self-administer testosterone orally (Johnson and Wood, 2001; Wood, 2002). In many studies of CPP, which is used to examine hedonic effects of drugs (Scoles and Siegel, 1986), T conditions a place preference (Alexander et al., 1994; Caldarone et al., 1996; DeBeun et al., 1992; Kashkin and Kleber, 1989; Packard et al., 1997, 1998; Schroeder and Packard, 2000), when administered systemically (Alexander et al., 1994) or when applied centrally to the NA (Packard et al., 1997) or to the medial preoptic area (King et al., 1999). However, there is considerable variability in this effect. In some studies, CPP with T was seen only with very high systemic dosages and not with lower dosages; in others, no effect was observed (Caldarone et al., 1996). The requirement for high dosages of T in order to produce effects on CPP is consistent with the notion that tolerance can be seen following repeated administration.

2.11. Tolerance and withdrawal

There is evidence from findings with people and animals that tolerance to AS can develop, which also leads to escalating and continued use. As early as 1950, Kochakian reported that the anabolic effect of AS were attenuated in rats repeatedly administered AS unless the dosages were increased. Up to 18% of AS users report tolerance (Brower et al., 1991). Withdrawal symptoms are also reported in rats given daily injections of T for 10, but not 3, weeks. For 2 weeks after T cessation rats had tremors, ataxic effects, and ptosis (Foltin, 1992). Although the findings discussed above indicate that androgens can produce interoceptive effects, what the nature of these effects is and the neurobiological substrates that mediate their actions remain to be elucidated.

3. Physiology and pharmacology of androgens

Steroids are vital for cell life. Early in evolution, hormones served as primitive growth regulators, and diversified later to sex steroids, gluco- and mineralo-corticoids with remarkable preservation of structure–activity relationships (Rousseau and
In mammals, all steroid hormones derive from cholesterol and are synthesized in steroidogenic organs, such as adrenals (mineralo- and gluco-corticoids), gonads and placenta (sex hormones including androgens), before being secreted into circulation. High lipophilicity of steroids facilitates penetration of biological membranes, securing access to all cells and organs, including the Central Nervous System (CNS).

The traditional view of how steroids exert their effects is through actions at specific intracellular steroid receptors. Briefly, once a steroid binds to its specific (cognate) intracellular steroid receptor, structural changes occur in the receptor that facilitate its binding to complementary regions of DNA in the cell nucleus. The receptor binding activates transcription of the gene(s), producing messenger RNA transcripts that encode a wide array of enzymatic, structural and receptor proteins (Rogozkin, 1991).

### 3.1. Genomic actions of androgens

Some of the effects of T and other AS may be mediated through intracellular androgen receptors (Janne et al., 1993). Androgen receptors are widely but selectively distributed throughout the brain (Stumpf and Sar, 1976). In the rat, the brain regions containing the highest levels of androgen receptors are the lateral septum, some areas in the hippocampus, the bed nucleus of the stria terminalis, the medial preoptic nucleus, the ventromedial hypothalamus, and the medial amygdaloid nucleus (Kritzer, 1997; Lieberburg et al., 1977). High dosages of AS lead to upregulation of androgen receptors in these areas (e.g., the ventral tegmental area, the CA-I region of the hippocampus), as well as in several non-classical target sites such as the locus ceruleus and the periaqueductal grey (Teledgy, 1987).

### 3.2. Non-genomic actions of androgens

Steroids may also influence cellular activity in a “non-genomic” fashion or through means other than traditional actions at intracellular steroid receptors (Brann et al., 1995). Indeed, T metabolites and many AS do not bind with a high affinity at cognate intracellular androgen receptors (Cunningham et al., 1979; Verhoeven et al., 1975). Some possible mechanisms for these non-classical actions are: changes in membrane fluidity; actions on receptors on plasma membranes; regulation of GBRs on plasma membranes; and, activation of steroid receptors by factors such as EGF, IGF-I and dopamine. These diverse intracellular and non-genomic modes of action provide for integrated actions of hormones which may be rapid and of short duration, or prolonged and of long duration.

There are well-preserved structure/activity relationships that are associated with genomic and non-genomic actions of steroid hormones. Hence, how a steroid works at a substrate may depend upon small differences in the steroids’ metabolism and structure. To a great extent, steroid effects may be determined by relatively few enzymatic steps performed by a small group of enzymes that result in the generation of all steroid hormones. In many respects, the metabolic pathway of a steroid determines the nature of the steroid signal and its degree of amplification (Rubinow and Schmidt, 1996). The final effect of a steroid hormone is determined by enzymatic activity in the target cell. For example, the composition of body hair is to a much higher extent related to 5α-reductase activity in the hair follicles than to plasma T levels (Lookingbill et al., 1991).

### 3.3. Androgen metabolism

Testosterone and many other androgens (i.e., those possessing a 3-keto 4-A configuration and a methyl group at the 19th carbon) may, via aromatization to estrogens (Akhtar et al., 1993; Graham-Lorence et al., 1995; Korzekwa et al., 1993, 1991), also stimulate estrogen receptors. Indeed, in situ aromatization of androgens represents an important metabolic event. Connolly et al. (1990) have demonstrated that adult male guinea pig brains contained higher quantities of androgen aromatase than female brains. Androgen aromatase is concentrated in the limbic system and hypothalamus (amygdala, preoptic area, septum, hippocampus, and medial basal hypothalamus), whereas low levels were consistently found in cortical tissue (McEwen, 1980). Furthermore, female rats treated with T show a more marked enhancement of social aggression than rats treated with the nonaromatizable androgen dihydrotestosterone (DHT). Combined treatment with DHT and estrogen resulted in the same degree of increased aggression as T treatment, suggesting that the activation of estrogen receptors and androgen receptors may work synergistically (Van de Poll et al., 1986). Many of the synthetic AS are aromatizable (Bahrke et al., 1990), a quality responsible for gynecomastia. Thus, central effects of AS may be mediated through genomic actions at androgen and estrogenic receptors. Furthermore, synthetic AS and their metabolites do not only bind to androgen or estrogen receptors but also to glucocorticoid and progestin receptors (Janne, 1990). Consequently, the effects of AS are far from purely androgenic and may involve actions at multiple genomic and non-genomic substrates.

Findings from our laboratory demonstrate that chronic administration of T and/or its 5α-reduced metabolites may have actions via both genomic and non-genomic substrates. Administration of 1 mg of T, DHT or its metabolite 3α-androstanediol (3α-diol) daily for 3 days reduced seminal vesicle weight (an androgen receptor dependent measure) and decreased androgen receptor binding in the hypothalamus. Concomitant with these androgen receptor mediated effects, the sensitivity of GBRs in the hippocampus was increased (Frye et al., 2001). As these androgen regimens are capable of having actions at both genomic (intracellular androgen receptors) and non-genomic (GBRs or other) substrates, an important question is what are the actions of androgens that underlie its hedonic effects.

### 4. Hypothesis

Our research to begin to address this question has focused on the role of T’s 5α-reduced metabolite, 3α-diol, which typically has actions at GBRs, rather than intracellular androgen receptors. The purpose of these studies is to better understand the abuse liability and potential of androgens’ by investigating the effects and mechanisms in mediating their interoceptive effects.
First, research is summarized that has investigated the extent to which T’s 5α-reduced metabolites may have actions in the nucleus accumbens (NA) to mediate hedonic effects. Second, experiments that address whether these effects of androgens require actions at intracellular androgen receptors is presented. Third, effects of facilitating and/or blocking androgens actions at GBRs in the NA is discussed. Finally, the results of effects of dopaminergic lesions to the NA on androgen’s effects are described. These findings that will be reviewed here suggest that 3α-diol’s actions in the NA via GBRs and/or dopaminergic neurons, rather than intracellular androgen receptors, may underlie hedonic effects (Fig. 1).

4.1. Are effects of T on conditioned place preference mediated by actions of its 5α-reduced metabolites?

Conditioned place preference (CPP) has been used in many experiments to examine hedonic effects of drugs of abuse (Scoles and Siegel, 1986). In several studies of CPP and androgens, T did condition a place preference (Packard et al., 1997; Alexander et al., 1994). However, there is considerable variability in this effect, with some studies reporting it only with extremely high dosages of T and not with lower dosages (Calderone et al., 1996), and in males but not in females (DeBeun et al., 1992). In some experiments rats were tested soon (30 min) after T administration, which may be insufficient time to enable T to be metabolized by 5α-reductase to DHT and by 3α-oxidoreductase to 3α-diol. As discussed below, the manner in which androgens are given, e.g., dosage, bioavailability, route of administration, and/or vehicle may underlie some of T’s variability on CPP.

Evidence from the literature to support the notion that variability in T’s positive hedonic properties may be related to capacity to form 3α-diol include the following. First, when administered systemically in oil vehicle 30 min prior to CPP chamber exposure only a high dosage (1 mg) of T was effective at inducing a CPP and lower dosages (10 or 100 μg) were not (Calderone et al., 1996). Higher dosages of T would more readily facilitate the metabolism of T even when administered in oil vehicle, compared to the lower dosages. Second, when administered systemically in a non-oil, encapsulization vehicle, 30 min prior to CPP chamber exposure, a CPP was observed in rats administered 1200 or 800, but not 400 μg/kg of T (Alexander et al., 1994). These dosages of T produced supra-physiological levels of circulating T in male rats (Taylor et al., 1989), suggesting that the rewarding affective properties of T depend upon circulating levels of hormones that are markedly above baseline concentrations. Third, when the bioavailability of T is further increased by intrabrain infusion of T in a molecular encapsulization vehicle, 0.25 or 0.50, but not 0.125 μg/kg of T into the NA of male rats immediately prior to CPP chamber exposure enhanced place preference (Packard et al., 1997). The metabolism enzymes, 5α-reductase and 3α-hydroxysteroid dehydrogenase, have been localized to the NA (Mellon, 1994) and could have rapidly converted the higher dosages of T to 3α-diol to produce these effects. The studies described below summarize our research progress to empirically address the question as to whether some of T’s hedonic effects may be related to formation of 3α-diol.

4.1.1. Systemic 3α-diol regimen that enhance CPP increase 3α-diol>DHT>T in plasma and NA

We have shown that systemic 3α-diol administration conditions a place preference more effectively than does systemic administration of DHT or T (Frye et al., 2001). Briefly, administration of

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Figure 1: Metabolism pathway and substrates for androgen action. Testosterone (T) is metabolized to dihydrotestosterone (DHT) via 5α-reductase, which is converted to 3α-androstanediol (3α-diol) via 3α-hydroxysteroid oxidoreductase. T and DHT bind to intracellular androgen receptors (ARs) and 3α-diol binds GBRs, which maybe localized to dopaminergic neurons in the nucleus accumbens.
3α-diol, but neither T nor DHT, to intact male Long–Evans rats, 1.0 mg daily for 6 days, 30 min prior to exposure to the non-preferred side of the CPP chamber, produced significant increases in preference for the non-preferred side of the chamber at testing (Fig. 2, right), compared to baseline preference (which did not differ across groups—Fig. 2, left). Notably, circulating concentrations of 3α-diol were increased most in 3α-diol>T-DHT>T-administered rats, compared to vehicle-administration (Fig. 2-inset).

Testosterone implants to the NA can condition a place preference (Packard et al., 1997); however, it is unclear whether this may be due to actions of its 5α-reduced metabolites. We have investigated effects of systemic androgens on CPP and levels of androgens in the NA (Rosellini et al., 2001; Frye et al., 2002). Rats were systemically administered 1 mg of T, DHT, or 3α-diol, 30, 90, or 180 min prior to exposure on conditioning days to the non-preferred side of a CPP chamber. All rats administered 3α-diol demonstrated CPP and had the highest concentrations of 3α-diol in the NA at each of the temporal pairings tested. The percentage of rats spending more time on the non-preferred side of the CPP chamber on the test day was greatest with androgen regimens that increased levels of 3α-diol in the NA. These findings demonstrate that 3α-diol concentrations are increased in the NA by androgen regimens that produce CPP (3α-diol>DHT>T-vehicle) (Rosellini et al., 2001).

4.1.2. Administration of T, DHT or 3α-diol directly to the shell of the NA enhances CPP
We have also investigated effects on CPP of directly stimulating the NA with androgens (Frye et al., 2002). Rats were administered implants of T, DHT or 3α-diol to the NA immediately prior to placement in the CPP apparatus on conditioning days. Implants of T, DHT, or 3α-diol immediately prior to exposure to the non-preferred side of chamber significantly increased time spent on the non-preferred side of the test day (Fig. 3, right), as compared to baseline (Fig. 3, left). Notably, this effect was only produced by androgenic stimulation of the shell, but not the core, of the NA (Fig. 3, inset). Results of this experiment and the latter suggest that androgen regimen that increase 3α-diol concentrations in the accumbens can enhance CPP and direct implants of T, DHT, or 3α-diol to the shell of the NA elicit CPP; however, whether formation of 3α-diol is required for these effects was not established. The data presented below, which heretofore have not been published, address the importance of androgen metabolism, actions at androgen receptors, GBRs, and dopamine targets.

4.1.3. Systemic metabolism inhibitors attenuates T-induced CPP and levels of 3α-diol in whole brain
Testosterone is metabolized by 5α-reductase to DHT; additional conversion by 3α-hydroxysteroid dehydrogenase forms 3α-diol. Findings above suggest systemic administration of 3α-diol>DHT>T paired with exposure to the non-preferred side of the CPP chamber produces the greatest effects on CPP and is associated with increases in 3α-diol in the NA during conditioning. These findings are consistent with the hypothesis that metabolism of T or DHT to 3α-diol in the NA is essential in mediating the hedonic effects of androgens. However, we have also directly tested the hypothesis that blocking T or DHT’s metabolism to 3α-diol will attenuate effects of the androgens on CPP. Systemic administration of 5α-reductase (finasteride) or by 3α-hydroxysteroid dehydrogenase (indomethacin) inhibitors, prior to T and/or DHT administration, attenuates CPP and whole brain 3α-diol levels.

Fig. 2. Rats administered subcutaneous T, DHT, or 3α-diol spend an increased amount of time on the non-preferred side of the conditioned place chamber and have increased levels of 3α-diol in the nucleus accumbens (inset). *P<0.05.

Fig. 3. Rats administered T, DHT, or 3α-diol to the shell of the nucleus accumbens spend an increased amount of time on the non-preferred side of the conditioned place chamber. 3α-diol administered to the core of the nucleus accumbens does not produce a place preference (i.e. no difference from vehicle administration to the core of the nucleus accumbens (inset). *P<0.05.
brain 3α-diol concentrations (Rosellini et al., 2003; Fig. 4, left). However, this does not address whether metabolism in the NA alone is required for hedonic effects of androgens.

4.1.4. Intra-accumbens metabolism inhibitors attenuates T-induced CPP and 3α-diol levels in NA

We have investigated whether androgen metabolism inhibitors to the NA can attenuate the hedonic effects of systemic T. First, finasteride applied to the NA prior to SC T paired with exposure to the non-preferred side of the chamber attenuates DHT and 3α-diol formation in the NA during conditioning trials as well and T’s facilitation of CPP at test time (Fig. 4, right). Second, indomethacin administered directly to the NA prior to SC DHT attenuates 3α-diol formation in the NA during conditioning and subsequently DHT’s facilitation of CPP at testing (Fig. 4, right). Third, the similar effects of systemic or intra-accumbens administration of metabolism inhibitors is very similar, which implies that blocking formation of 3α-diol in the NA is sufficient to attenuate T’s effects on CPP. Fourth, similar effects of blocking T’s and/or DHT’s metabolism suggest that 3α-diol, rather than DHT, in the accumbens is the active androgen underlying some effects on CPP.

4.2. Are actions at androgen receptors in the NA necessary for androgens effects on CPP?

The findings described above that 3α-diol and its prohormones T and DHT can induce CPP when applied to the NA, a region of the brain with few intracellular androgen receptors (Stumpf and Sar, 1976), suggests that these androgens may exert some of their hedonic effects via non-genomic actions. For example, implants of T to the medial preoptic area or the NA condition a place preference (Packard et al., 1997). As well, implants of T or 3α-diol also condition a place preference when applied to the NA (Frye et al., 2002; Rosellini et al., 2001).

Although there are many intracellular androgen receptors in the medial preoptic area that could be substrates for T’s effects, there are few intracellular androgen receptors in the NA (Stumpf and Sar, 1976). As well, we have shown that implants of T or 3α-diol and immediate pairing with the non-preferred side of the chamber enhances CPP. Testosterone and DHT both bind readily to intracellular androgen receptors, while 3α-diol is devoid of affinity for androgen receptors in physiological concentrations (Cunningham et al., 1979; Verhoeven et al., 1975). Although there are few androgen receptors that have been identified in the NA and the rapid effects of 3α-diol to enhance CPP would seem to preclude sufficient time for androgen receptor mediated changes in transcription, it is necessary to investigate whether 3α-diol’s actions at for androgen receptors are necessary for its hedonic effects. Pharmacological concentrations of 3α-diol that enhance CPP could override 3α-diol’s selective low affinity for androgen receptors and thereby potentially produce effects via the few androgen receptors in the NA or in other regions of the brain, such as the hippocampus.

4.2.1. Systemic administration of an androgen receptor blocker does not attenuate 3α-diol-induced CPP

We investigated whether 3α-diol has actions via intracellular androgen receptors to mediate CPP. During conditioning, rats received SC flutamide (10 mg) or vehicle (sesame oil) 2 h prior to SC 3α-diol (10 mg/kg) or vehicle (propylene glycol), which were administered immediately before placement in the non-preferred side of the CPP chamber. As we have previously demonstrated, rats that received vehicle and 3α-diol spent a significantly increased amount of time on the non-preferred side of the chamber at test time compared to vehicle–vehicle or vehicle–flutamide controls. Co-administration of flutamide with 3α-diol does not attenuate 3α-diol-induced CPP, and did not have intrinsic effects (Fig. 5, left). Although, these findings suggest that 3α-diol’s actions to enhance CPP occur independent of actions at intracellular androgen receptors in any part of the brain, the flutamide regimen utilized has been demonstrated to block physiological, rather than pharmacological regimen of androgens. As such, we also wanted to investigate effects of a pharmacological flutamide regimen to counter pharmacological effects of 3α-diol.

4.2.2. Intra-accumbens administration of an androgen receptor blocker does not attenuate T-induced CPP and 3α-diol levels in NA

We investigated whether 3α-diol has actions via intracellular androgen receptors in the NA to mediate CPP. Rats were administered flutamide or empty control implants to the NA 2 h prior to SC 3α-diol or vehicle immediately prior to placement in the non-preferred side of the CPP chamber. As previously demonstrated, rats administered 3α-diol exhibited a place preference for the non-preferred side of the chamber on test day compared to vehicle. Administration of flutamide implants directly to the NA with 3α-diol did not attenuate 3α-diol-induced CPP, nor did it effect CPP itself in the absence of 3α-diol (Fig. 5, right). Although these data indicate that actions of 3α-diol in the NA to mediate CPP may be independent of its...
actions at ARs, the neurobiological substrates in the NA that mediate 3α-diol’s hedonic effects need to be revealed.

4.3. Are actions at GBRs in the NA necessary for androgens effects on CPP?

There are data to suggest that steroids also may influence cellular activity in a “non-genomic” fashion or through means other than traditional actions at intracellular steroid receptors (Brann et al., 1995). Indeed, T, its metabolites, and many AS, do not bind with a high affinity to traditional intracellular androgen receptors (Cunningham et al., 1979; Verhoeven et al., 1975). Anabolic steroids upregulate androgenic substrates in several non-classical target sites, e.g., the CA-I region of the hippocampus and the VTA (Teledgy, 1987). Likely mechanisms for these non-classical actions are: (a) changes in membrane fluidity; (b) steroid hormones acting on receptors on plasma membranes; (c) steroid hormones regulating GBRs on plasma membranes; and (d) activation of dopamine substrates. The GABA system has been implicated as a substrate for illicit drug use, and steroids have been shown to interact with GBRs (Majewska et al., 1986; Gee, 1988; Frye et al., 1996b) and dopamine systems (Mani et al., 1996; Frye et al., 2000). Although the former (a and b) possibilities cannot be ruled out, we have not investigated these possibilities to date because such substrates have not been sufficiently defined as to warrant extensive investigation at this time. Our research to begin to address the possibility that GBRs (this section) or dopaminergic neurons (d; next section) are possible “non-genomic” substrates for androgens’ hedonic effects is described below.

Evidence for GABA involvement in AS mechanism of action include that withdrawal is characteristic of drugs of abuse that act at GABA. Withdrawal from chronic exposure to other psychoactive GABA-active agents, such as ethanol, benzodiazepines, and barbiturates, can result in psychomotor disruption and anxiogenic effects (Buck et al., 1991; Hauser et al., 1989; McCaslin and Morgan, 1988). Cessation or diminished use of AS has been associated with depressive symptoms (Corrigan, 1996). Other symptoms related to the loss of positive psychological effects of AS include listlessness; apathy; loss of appetite, libido and self-esteem; feelings of anxiety; difficulty in concentrating; and mood swings (Bahrke et al., 1990; Corrigan, 1996; Uzych, 1992).

There is evidence that androgens, particularly 3α-diol, may have actions through GBRs. Several steroids can alter GBR function, and the 5α-reduced, 3α-hydroxylated structure of 3α-diol meets the requirements of the most potent steroid modulators of GBRs (Belelli et al., 1990; Beyer et al., 1988). Although T is not particularly effective at altering GBRs, 3α-diol is (Gee, 1988; Frye et al., 1996a) and GBRs have been localized to the NA (Zhang et al., 1991). Chronic administration of T and metabolites alters the sensitivity of GBRs in the hippocampus (Frye et al., 2001). AS administration stabilizes GBRs in a moderate affinity state for benzodiazepine binding and reduces the EC50 for GABA-stimulated chloride influx (Masonis and McCarthy, 1995, 1996). GABA-stimulated chloride influx and muscimol binding are increased in animals administered 3α-diol (Frye et al., 1996a,b,d). Co-administration of a GBR antagonist, bicuculline, counters 3α-diol-induced changes in social (Frye et al., 1996a), and affective (Frye et al., 1996c) behavior. Furthermore, rats administered the AS, dianabol, demonstrated increases in levels of 3α-diol in the NA and greater GABA-stimulated chloride influx in cortical synaptoneurosomes than did vehicle controls or rats that had dianabol discontinued and were withdrawing (see Table 1).

4.3.1. Intra-accumbens GBR antagonist attenuates 3α-diol-induced CPP

These background data that show that 3α-diol has actions via GBRs suggest that androgens’ hedonic effects may be mediated in part via GBRs in the NA. To further determine the role of GBRs in the NA to mediate some of the effects of androgens on CPP, the following experiments were conducted. Rats received either a high dosage of 3α-diol (1 mg) that has been used previously to induce CPP or vehicle priming, in conjunction with infusions of vehicle or the GBR antagonist, bicuculline (50 ng) to the NA immediately prior to exposure to the conditioning chamber. As previously demonstrated, rats administered 1 mg of systemic 3α-diol showed an increase in time spent on the non-preferred side of the

<table>
<thead>
<tr>
<th>Treatment group</th>
<th>3α-Diol levels (ng/ml)</th>
<th>Emax</th>
<th>EC50 (ng/ml)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Vehicle</td>
<td>2.25±0.5</td>
<td>24.4±1.4</td>
<td>11.2±0.7</td>
</tr>
<tr>
<td>Dianabol</td>
<td>3.00±0.2</td>
<td>34.8±3.3 *</td>
<td>9.3±0.7 *</td>
</tr>
<tr>
<td>Dianabol</td>
<td>1.40±0.2</td>
<td>28.1±4.0</td>
<td>11.8±1.0</td>
</tr>
<tr>
<td>Withdrawal</td>
<td></td>
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* Indicates a significant (P<0.05) difference from vehicle-administered controls.
conditioning chamber on test day compared to rats that received vehicle. Co-administration of bicuculline, a GBR antagonist, to the NA, attenuated 3α-diol-induced CPP. Neither bicuculline administration alone, nor vehicle administration alone, had effects on CPP (Fig. 6). Although these findings suggest that blocking 3α-diol’s actions at GBRs can attenuate CPP; a truer measure of whether 3α-diol can work through GBRs to produce hedonic effects would involve facilitation of such effects.

4.3.2. Intra-accumbens GBR agonist enhances 3α-diol-induced CPP

In this experiment, we examined whether activating GBRs could enhance effects of subthreshold 3α-diol stimulation to produce CPP. Rats were systemically administered a lower dosage of 3α-diol (1 mg/kg) that alone is insufficient to induce CPP or vehicle-priming. In addition, the GBR agonist, muscimol, or saline vehicle was infused to the NA immediately prior to being put in the conditioning chamber. As expected, rats administered the lower dosage of 3α-diol (1 mg/kg) spend a similar amount of time on the non-preferred side of the chamber on test day as did vehicle-primed rats. However, co-administration of muscimol, a GBR agonist, to the NA together with this sub-threshold dosage of 3α-diol, was sufficient to induce a CPP. Muscimol alone did not produce a CPP and was no more effective than vehicle at enhancing CPP (Fig. 7). These data, which show that 3α-diol and muscimol may have synergistic actions as GBR agonists, suggest that 3α-diol may have effects in the NA on CPP in part through agonist-like actions at GBRs.

4.4. Are actions of 3α-diol at dopaminergic neurons essential for 3α-diol-enhanced CPP?

Interceptive effects of androgens may involve direct or indirect actions at dopaminergic neurons. Evidence in support of this is as follows. We, and others (Packard et al., 1997), have found that androgens can condition a place preference when applied directly to the NA, an area of the brain containing dopamine neurons. Administration of AS can produce EEG activity similar to that of psychostimulants (Itil, 1976; Itil et al., 1974; Stenn et al., 1972). Androgens can act on the dopamine reward system in a manner similar to cocaine or other stimulants (Alderson and Baum, 1981; Goudsmit et al., 1990; Jalilian-Tehrani et al., 1982; Mitchell and Stewart, 1989; Kashkin and Kleber, 1989; Vermes et al., 1979). Notably, the mesolimbic dopamine system is often considered the final common pathway for many dependence-producing drugs (Koob and LeMoal, 1997; Koob, 1992; Wise and Bozarth, 1987; Robinson and Berridge, 1993). There are some reports of AS use enhancing sexual desire and pleasure in people (Greenblatt and Karpas, 1983; Taylor, 1987), effects that are known in rats to be associated with increased levels of dopamine in the NA (Moses et al., 1995; Lorrian et al., 1999). We have begun to address whether another possible non-genomic substrate through which androgens may have their hedonic effects is via dopaminergic neurons in the NA.

4.4.1. 6-OHDA lesions that completely eradicate the shell of the NA prevent 3α-diol’s enhancement of CPP

In this study, we hypothesized that 6-hydroxydopamine (6-OHDA) lesions to the dopamine neurons of the NA would decrease the conditioning effects normally seen with 3α-diol administration. 3α-diol was administered subcutaneously (SC) 1 mg daily for the...
6 days of pairings with the non-preferred side of the chamber to 6-OHDA lesioned male Long–Evans rats, 30 min prior to exposure to the non-preferred side of the CPP chamber. 3α-diol conditioned a place preference in rats with partial lesions to the shell and core of the NA, but no conditioning was seen in rats that had complete lesions to the shell of the NA (Fig. 8). These data suggest 3α-diol’s enhancing effects on CPP may require actions at dopaminergic neurons in the shell of the NA. This suggestion is consistent with the conclusion reached by McBride et al. (1999) in their thorough review of the literature on brain reinforcement mechanisms that intracranial self-administration and conditioned place preference studies with “psychostimulants, morphine, and PCP all produce reinforcing effects within the NA and that their (reinforcing) effects are observed mainly in the shell” (p 141). As such, these data, and the findings discussed above that 3α-diol implants to the shell, but not the core, of the NA enhance CPP, also underscore other findings presented, which indicated that 3α-diol can have positive hedonic effects and that such effects may involve actions at GBRs and dopamine neurons.

5. Other substrates for AS

This review of potential targets for androgens’ actions for hedonic effects is by no means exhaustive. There are clearly numerous substrates at which androgens and AS could have actions to produce positive hedonic effects. One that may be particularly relevant that is beyond the scope of the present review, but warrants further consideration, is the opioid system. Many dopamine neurons in the mesolimbic pathways are juxtaposed with opioid containing neurons. Among the reported interoceptive effects of AS in people are effects similar to that seen in narcotic users, such as euphoric mood (Pope and Katz, 1988, 1994) and elevation of mood in depressed patients (Bahrke et al., 1990). Indeed, there is evidence that AS use may be increasingly considered a “gateway” drug for later use of opiates (McBride et al., 1996; Pope et al., 2000). In 1999, 10% of men being treated for opiate addiction in a rehabilitation program indicated that AS use preceded their opiate use. In 1990, only 1% of men in treatment for dependence in the same rehabilitation program reported any preceding AS use (Arvary and Pope, 2000). There is also evidence that withdrawal from AS may share many common characteristics of opiate withdrawal, such as the initial phase of withdrawal characterized by increased pulse and blood pressure, sweats, chills, nausea, and dizziness (Kashkin and Kleber, 1989; Tennant et al., 1988).

6. Summary

In summary, androgen and AS use is widespread and increasing. The background literature regarding effects of AS underscore the notion that AS can have effects that are akin to that of other drugs of abuse. Use of AS is associated with adverse illicit use of other drugs of abuse, as well as physiological and behavioral consequences (including violence and aggression). Anabolic–androgenic steroids use may lead to addiction, dependence, and withdrawal such that use is often continued despite short-and long-term health risks. Understanding the pharmacological effects of AS is important, but attempts have been confounded both by the complex receptor-and non-receptor-bound actions of AS and the lack of basic understanding of androgens’ actions and their hedonic effects and neurobiological mechanisms. The research described addresses whether T, a widely used AS, has positive euphorogenic effects which may contribute to their abuse liability in part through non-genomic actions. The implications of this research are that AS and/or T may produce some of their positive hedonic effects by enhancing 3α-diol production, which in turn has actions at GBRs in the NA, which synapse on dopaminergic neurons, to produce positive hedonic effects. Further research is needed to ascertain other hedonic effects and mechanisms of androgens.

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