

Canadian Urological Association guideline on the care of the normal foreskin and neonatal circumcision in Canadian infants (full version)

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Introduction

Circumcision is the oldest planned operative procedure in the history of the human civilization, but there continues to be a lack of consensus and strong opposing views on whether universal neonatal circumcision should be adopted as a public health measure. The recent American Academy of Pediatrics (AAP) guideline on male circumcision (MC) reversed its prior stand, stating that the “health benefits of newborn male circumcision outweigh the risks,” and justify access to the procedure if the parents so choose.¹ This recommendation was primarily based on the impressive results from African trials demonstrating the protective effect of MC against human immunodeficiency virus (HIV) and sexually transmitted infections (STIs).

A review of the literature on MC shows evidence of a vehement debate, often clouded by strong personal biases and lack of high-quality evidence. Creation of a guideline specific to the need of Canadian infant males is, therefore, difficult given the level of evidence provided for each potential benefit, the lack of data directly applicable to the Canadian population, the inability to quantify the true complication rate of routine circumcisions accurately, uncertainty about the health benefits of a circumcision compared with other health interventions, the ethical issues and acceptability of a surgical procedure done by parental consent for future benefits, and the costs of training and implementation of any universal neonatal circumcision policy in Canada.

Acceptance and trends of neonatal circumcision

Neonatal circumcision rates are declining across several countries, including Canada, and this may be a reflection of changing demographic patterns and parental beliefs. The

Provincial Ministries of Health in Canada indicate circumcision rates of 51–67% in 1970. In 2009, the Public Health Agency of Canada reported an overall Canadian circumcision rate of 31.9% for 2006–2007. This varied across the provinces, with the rate being highest in Alberta (44.3%) and lowest in Nova Scotia (6.8%) (www.circinfo.net).

A Centers for Disease Control and Prevention (CDC) report showed a decreasing trend in US circumcision rates from 1999–2000 until 2008–2010 from 60% to 55% of newborn males.² In the U.K. between 1997 and 2004, circumcision rates declined from 2.6/1000 boys/year to 2.1/1000 boys/year;³ however, a more recent study from the U.S. using the Nationwide Inpatient Sample noted an increasing trend of neonatal circumcisions from 48% in 1988–1991 to 61% in 1997–2000, reflecting an increase of 6.8% on average every year.⁴

A survey of prospective parents in the U.S. was assessed to obtain parental views of circumcision analyzing the effect of the AAP 1999 circumcision guideline and recent HIV and human papillomavirus (HPV) trials.⁵ Individuals with previous circumcised sons, those born in the U.S., and those who discussed circumcision with their partner were more likely to request circumcision. There was no change in support after reading information on recent HIV/HPV trials from Africa. A similar survey in Canada indicated that circumcision status of the father significantly influences the child's circumcision.⁶ This study indicates that initial parental views rather than new evidence may continue to be the strongest determinant for neonatal circumcision.

Aims of the review

The aim of this guideline is to present the current evidence on the benefits of circumcision, the optimal anesthesia/analgesia requirements of neonatal circumcision, the possible complications of circumcision, and its effect on sexual function and sensation, as well as the care of a normal foreskin in early childhood. This guideline is directed towards pediatric caregivers who routinely examine and follow male infants,

physicians who provide neonatal circumcision services, and pediatric urologists and general surgeons. The current guideline is written with the purpose of being applicable to the Canadian population and healthcare system. The evidence presented is classified according to the Oxford system of evidence-based medicine.⁷

The current guideline attempts to answer the following questions:

1. Do the potential benefits of neonatal circumcision justify performing universal neonatal circumcision in Canada?
2. For an individual patient and parent, what are the benefits and risks of a neonatal circumcision and how reliable and applicable is the evidence currently available?
3. What should be the prescribed routine foreskin care in infants, indications for medically indicated circumcision, and management of physiological phimosis?

Methods

Systematic literature searches were conducted in MEDLINE, including Pre-MEDLINE EMBASE, BIOSIS Previews®, Web of Science® — with conference proceedings, and the Cochrane Central Register of Controlled Trials electronic bibliographic databases, and were restricted to either adult or pediatric studies (</> 18 years) (2002 to March 2013). All searches were restricted to studies published in English language only.

The search queries were developed using a combination of subject headings and free-text terms such as circumcision, circumcision male, uncircumcised, male sexual dysfunction, sexual dysfunction physiological, sexual dysfunctions psychological, erectile dysfunction, sexual problems, sexual arousal disorder, ejaculation dysfunction, sexuality, prostatic neoplasms, prostate cancer, prostate tumour, penile neoplasms, penile cancer, urinary tract infections, phimosis, HIV infections, HIV, human immunodeficiency virus, HPV infections, STIs, using variant spellings and endings. For all searches, editorials, news, and letters were excluded. The bibliographies of all relevant retrieved articles and reviews were also examined to identify further relevant articles.

A total of 2674 records were identified and after removing duplicate records and excluding non-relevant studies, 229 studies were identified for detailed analysis and included in this analysis.

Care of the normal foreskin in childhood and management of physiological phimosis

Natural history of the foreskin

The prepuce arises from the coronal margin by a combination of folding and epithelial outgrowth and has an outer and inner layer separated by Dartos fascia. At birth, the inner foreskin is usually fused to the glans penis and should not be forcibly retracted unless it is possible to retract it with gentle traction. Initial examination of the newborn with a normal prepuce without any ventral deficiency or dorsal hood is usually (except in a baby with a megameatus intact prepuce variant of hypospadias) a reliable indicator that the urethral meatus is in a normal location and rules out significant hypospadias.

The collection of smegma (a white exudate of skin cells and keratin) separating the prepuce from the glans and repeated reflex erections are the primary mechanisms that lead to resolution of physiological adhesions over time. This process is usually complete by three years of age in 90% of the boys, although this study is a reflection of the poor current data in this regard.⁸ In a more recent study from Taiwan, the incidence of non-retractable physiological phimosis was 50% in grade 1 boys and decreased to 35% in grade 4 and 8% in grade 7 boys.⁹

When does physiological phimosis require treatment?

In the absence of clinical findings of scarring suggesting pathological phimosis and history of recurrent urinary tract infections (UTIs) or balanoposthitis, no intervention is required for physiological phimosis. Ballooning of the foreskin during voiding is not associated with obstructed voiding and is not an indication for circumcision.¹⁰ Active retraction has the potential to cause microtears and lead to scarring and subsequently a true phimosis. Therefore, normal foreskin care in early childhood only starts once the foreskin is retractable and this will happen at varying ages.¹¹ Once retractable, the child can be taught to gently retract and clean during bathing with normal soap and water.

Indications for urological consultation in this age group include suspicion of true phimosis with evident scarring of the preputial ring (Fig. 1), recurrent episodes of balanitis (Fig. 2), genital lichen sclerosis (Fig. 3), or UTI.⁴ If the foreskin is not open by 8–10 years of age, there may be an indication for steroid therapy and gentle retraction, though there is no prescribed, evidence-proven age cutoff for this process.



Fig 1. Pathological phimosis.

Treatment of physiological phimosis

Several observational studies and randomized trials have investigated the role of topical steroids and preputial stretching in resolving physiological phimosis. The key to success with these protocols lies in differentiating physiological and true phimosis, active counselling and patient selection. Topical steroid aids by thinning the preputial skin and obliterating the stratum corneum, which then allows gentle retraction over time.

Level 1 evidence

Letendre conducted a randomized, double-blind study comparing a two-month treatment course of emollient cream vs. 0.1% triamcinolone in boys 3–12 years of age.¹² At four months, 76% of those on triamcinolone cream responded as compared to 39% in the placebo group ($p=0.008$), with no complications. At one year followup, the success rate in the steroid group had come down to 47%. In another randomized controlled trial (RCT) comparing a moderately potent steroid mometasone to moisturizing cream, the authors found a significantly better response rate of 88% at eight weeks compared to 52% in the placebo group.¹³ In a randomized, double-blinded, placebo-controlled study using betamethasone, Lund et al showed an initial 74% cure rate at four weeks' followup.¹⁴ At 18 months, 14% showed a relapse, but none required a circumcision. A lower success rate of 52% was noted by Nobre et al in a RCT from Brazil using 0.2% betamethasone-hyaluronidase cream in boys 3–10 years of age.¹⁵

Success rates did not vary by steroid potency, as shown by similar foreskin retraction rates using clobetasone (moderately potent steroid, success rate 77%) compared to betamethasone (highly potent steroid, success rate 81%).¹⁶ Side effects



Fig 2. Balano-posthitis.

are rare and there was no suppression of the hypothalamic-pituitary-adrenal axis provided treatment is not prolonged beyond eight weeks for each course.¹⁷

Level 2–4 evidence

Zavras et al conducted a prospective study using a mildly potent steroid, fluticasone propionate (0.05%), to achieve a 91% success rate in 1185 boys referred with a diagnosis of phimosis.¹⁸ Long-term success is maintained in over 75% of the boys following initial success with steroid therapy.^{19,20} Ku et al noted that success rates were higher in boys <3 years of age (92%) compared to those ≥ 3 years (70%), which may



Fig 3. Lichen sclerosus of the foreskin.

reflect compliance or a higher likelihood of pathological phimosis in the older group.²¹ Ashfield et al showed a reverse trend of better success rate in older boys, although the result was not found to be statistically significant at any age cutoff.²² Elmore et al showed that topical steroids as an alternative to circumcision are equally effective (74%) and safe in infants presenting with genitourinary abnormalities and UTIs.²³

Recommendations (care of the normal foreskin and physiological phimosis):

- 1. Neonatal examination of the foreskin and urethral meatus should be part of routine clinical assessment of all newborn boys. Continued examination of the foreskin without forcible retraction is recommended during yearly physical examinations to rule out pathological phimosis and document natural preputial retraction (Level 5, Grade D).**
- 2. Persistent physiological phimosis in an asymptomatic child should not be an indication for circumcision (Level 5, Grade D).**
- 3. Physiological phimosis requires treatment if associated with true balanoposthitis or recurrent UTIs (Level 5, Grade D).**
- 4. Topical steroids are the first-line treatment for persistent physiological phimosis requiring treatment with good success rates and low risk of complications (Level 1b/2b, Grade A).**
- 5. Moderately low-potency steroid (triamcinolone, clobetasone, hydrocortisone, mometasone) may have similar success compared to a highly potent steroid (betamethasone) (Level 2b, Grade B).**
- 6. Patient selection to ensure compliance, demonstrating the technique of gentle retraction of the foreskin and continued retraction after initial success is important to achieve continued success to topical steroid therapy (Level 5 Grade D).**
- 7. Recurrence of physiological phimosis is common and normally responds to another course of topical steroids (Level 2b/3 Grade C).**

Circumcision and risk of UTI

Prior evidence indicates that neonatal circumcision decreases the risk of UTI. The current debate centres on the magnitude of this effect, the overall effect given the low prevalence of male UTI, the lack of high-level evidence, and the need for a surgical procedure to prevent this risk. The role of circumcision in preventing UTIs must be studied in two distinct subgroups: males with normal urinary tracts and those with recurrent UTIs or urological conditions predisposing to UTI, like vesicoureteric reflux, posterior urethral valves, neurogenic bladders, and primary megaureters.

Childhood UTI epidemiology

Prior data suggest that in boys without predisposing urological conditions, the estimated incidence of UTI in the first 10 years of life varies from 1–2%.^{24,25} The prevalence rate of UTI in symptomatic children is higher. In a meta-analysis of 18 studies, Shaikh et al estimated that in all febrile infants (males and females 0–24 months), the prevalence of UTI was 7% (95% confidence interval [CI] 5.5–8.4).²⁶ In older symptomatic children (2–19 years), the prevalence was 7.8% (95% CI 6.6–8.9). Males under three months of age had the highest prevalence of UTI (8.7%; 95% CI 5.4–11.9). There was evidence of significant heterogeneity, but no publication bias and bagged specimens were included in some studies.

Biological plausibility for the role of circumcision in UTI prevention

Circumcision prevents UTI by reducing periurethral bacterial colonization secondary to reduced adherence of bacteria to keratinized surfaces and by removing the growth-promoting moist preputial environment.^{27–29} Foreskin colonization with potential pathogenic bacteria occurs early and leads to an increase in protective Langerhans cells.³⁰ In a case-control study, circumcision decreased the bacterial colonization of the glans penis for uropathogenic bacteria when compared to boys with an intact foreskin and this effect persisted in older boys.³¹ Studies have shown *E. coli* strains causing UTI in uncircumcised male infants resemble urosepsis strains isolated in adults.³² In addition, data from the HIV type 1 trial in Uganda showed a decreased prevalence and load of 12 specific anaerobic bacteria following circumcision.²⁷

Boys with normal urinary tracts

Level 2 evidence

In a meta-analysis, among febrile male infants less than three months of age, the prevalence of UTI was 10 times higher in uncircumcised males (20.1%; 95% CI 16.8–23.4) than circumcised males (2.4%; 95% CI 1.4–3.5).²⁶ This difference decreased in the 6–12-month group (7.3% and 0.3%) and there was no data available beyond infancy. Translated to likelihood ratios (LR), the uncircumcised male infant (3–24 months old) has a UTI LR of 2.8 vs. 0.33 for circumcised males.³³ The risk increases if this infant is of non-black race. Singh-Grewal performed a meta-analysis of 12 published studies (one RCT, four cohort, seven case-control) on 402 908 children published until 2002 and, assuming a 1% risk of UTI, calculated the number of circumcisions required to prevent one UTI as 111.³⁴ The quality of the included studies was poor, with variable UTI definitions, bagged specimens in at least six studies, and potential differential misclassifi-

cation, as 3/4 of the cohort studies were unable to account for circumcisions past the neonatal period. The overall odds ratio (OR) for a UTI in circumcised males compared to uncircumcised males was 0.13 (95% CI 0.07–0.23). The single RCT by Nayir et al included in this analysis compared bacteriuria rates between circumcised and uncircumcised boys using bag or mid-stream specimens and showed a non-statistically significant OR of 0.13 in the circumcised group (95% CI 0.01–2.63).³⁵

A systematic review conducted by Morris et al, calculated the lifetime risk of a UTI to be 32% in uncircumcised males compared to 9% in circumcised males.³⁶ The authors suggested a number needed to treat (NNT) of 4.2 (95% CI 2.2–27) for preventing one UTI over a lifetime. A Cochrane review in 2012 failed to identify any new RCTs to include in a meta-analysis.³⁷

Level 2–4 evidence

Zorc et al conducted a prospective cross-sectional study to identify clinical factors associated with UTI in infants <60 days of age.³⁸ The overall rate of UTI was 9% and after multivariable adjustment, being uncircumcised was associated with a higher UTI risk (OR 10.4; 95% CI 4.7–31.4; $p < 0.001$). According to a large population-based cohort study conducted in Canada by To et al, the relative risk of UTI requiring hospital admission in uncircumcised compared to circumcised boys was 3.7 (95% CI 2.8–4.9) in the first year of life.³⁹ The number of circumcisions needed to prevent one admission for UTI in the first year of life was 195. In another well-conducted, case-control study from Australia, which included children with urological abnormalities, Craig et al showed that the protective effects of circumcision in reducing risk of UTI extended beyond infancy.⁴⁰ In infants, the OR was marginally significant at 0.03 (95% CI 0.06–1.1); however, the OR for the >1-year group was not statistically significant (OR 0.2; 95% CI 0.01–3.7).

The rate of asymptomatic positive urine cultures (bag specimens confirmed with suprapubic aspiration) was significantly lower in a study by Simforoosh et al.⁴¹ None of the 3000 circumcised children followed up to 15 months age had a positive culture, while 2% of the 1000 uncircumcised males had a positive culture. Interestingly, several studies from Israel, where all males undergo a ritual neonatal circumcision, showed a high incidence of post-circumcision UTI.^{42,43}

Boys with abnormal urinary tract

In urological conditions, like high-grade vesicouretric reflux (VUR), posterior urethral valves, and primary megaureters, the risk of UTI is higher.

VUR

In a prospective cohort study, Alsaywid et al noted a lower non-significant incidence of new defects on dimercaptosuccinic acid (DMSA) scans in circumcised males with Grade 4–5 VUR as compared to uncircumcised boys (5.25 vs. 10.2%);⁴⁴ however, three of the four boys in the circumcised group with new DMSA defects did not get breakthrough UTIs. Circumcision was more effective than antibiotic prophylaxis alone or anti-reflux surgery in preventing breakthrough UTI (OR 0.9). In contrast, after surgical correction of VUR, a concomitant circumcision did not decrease the risk of post-operative UTI.⁴⁵ In a small RCT conducted on children <3 years with low-grade VUR (Grade 1–3) randomized to antibiotic prophylaxis and prophylaxis plus circumcision groups, the authors noted significantly lower positive peri-urethral cultures in the circumcised group up until nine months of followup, following which, results equalized.⁴⁶ The authors also showed a significantly lower positive culture rate by urethral catheterization in the circumcised group, but did not comment these patients were symptomatic. Posterior urethral valves

Mukherjee et al showed, in a retrospective cross-sectional study, that circumcision in boys with posterior urethral valves significantly reduces the incidence of UTI beyond infancy by 83%.⁴⁷ Given that their cohort had a very high incidence of UTI, the NNT to prevent one UTI was one in boys with valves.

Ureteropelvic junction (UPJ) obstruction and antenatal hydronephrosis

In a recent prospective study on infants with antenatally detected hydronephrosis, uncircumcised boys (adjusted OR 3.63; 95% CI 1.2–11.2) and females had a significantly higher risk of febrile UTI compared to circumcised males.⁴⁸

Roth et al did not demonstrate a protective effect of circumcision in children with Grade 3–4 hydronephrosis secondary to UPJ obstruction or obstructive megaureters in a retrospective cohort study.⁴⁹ Although unable to show statistical significance, the UTI rate was 0 in circumcised males as opposed to 8.3% in uncircumcised males. In addition, there is indirect evidence of the protective effect of circumcision in this population when comparing this study (63% circumcised), which had a 4.3% UTI rate, with a similar cohort presented in a study by Song et al, which had a higher 36% UTI rate (0% circumcised).⁵⁰

It is unclear whether medical treatment of physiological phimosis or antibiotic prophylaxis can prevent UTIs as effectively as circumcision in this subset of males. Urine specimen collection methodology is an issue and a recent study demonstrated that even catheterized specimens in uncircumcised boys could be contaminated.⁵¹ Studies measuring the incidence of UTIs are intrinsically more difficult to interpret, as most measure number of UTI episodes as opposed to

number of children with UTI. Assuming a 1–2% circumcision complication rate and a 1% UTI risk in normal infants, universal neonatal circumcision cannot be justified based on a NNT of 111.³⁴ Even if we accept a lower complication rate of 0.2% and a 2% UTI risk, given the effectiveness of UTI treatment, six UTIs will be prevented at the expense of one circumcision complication. This equation changes in favour of a circumcision in those with recurrent UTI (assuming a risk of 10%, the NNT is 11) and boys with urological abnormalities (even assuming an inflated risk of UTI with VUR being 30%, the NNT is 4).³⁴

Recommendations (MC and UTI prevention):

- 1. Neonatal circumcision decreases the risk of UTI (Level 2a).**
- 2. The risk of UTI is low in infant males and decreases further beyond infancy (Level 2b–4).**
- 3. There is paucity of Level 1 evidence to justify recommending universal circumcision to prevent UTIs in normal males.**
- 4. A stronger effect of neonatal circumcision in preventing UTIs in boys with urological abnormalities has been demonstrated and, therefore, it is recommended that a discussion with the parents is advisable for this subgroup of neonates (Level 3–4 Grade C).**

Circumcision and risk of sexually transmitted infections

A decreased risk of HIV and other STIs is the primary argument driving a change in risk-benefit assessment of MC. The other benefits of MC are overshadowed to a large extent by its effect on HIV and STIs and, therefore, data in relation to these benefits must be scrutinized carefully to obtain evidence-based recommendations.

1. Circumcision and HIV prevention

Epidemiology of HIV infections

A recent Public Health Agency of Canada report estimated a HIV prevalence rate of 208 per 100 000 population, with an 11.4% increase compared to 2008 estimates.⁵² Men who have sex with men (MSM) accounted for 47% of prevalent infections, followed by intravenous drug users (17%) and heterosexual individuals (17.6%).

Based on RCTs conducted in HIV high prevalence sub-Saharan Africa, there is clear Level 1 evidence that MC reduces the risk of HIV infection in heterosexual men in that region of the world.^{53–55} In 2007, the WHO and UNAIDS recommended MC, regardless of HIV serostatus, as an additional intervention in countries with predominant heterosexually acquired HIV infection, where HIV is prevalent and circumcision rates are low.⁵⁶ Circumcision should not

replace promotion of the “ABC” (abstinence, behaviour change, and correct and consistent condom use) strategy or value of voluntary counselling and testing and care services for STIs. The CDC convened a consultation on public health issues regarding male circumcision in the U.S. for prevention of HIV infection in 2007 and put forward the following key proposals:⁵⁷

1. With respect to HIV prevention, MC is one of several partially effective risk-reduction alternatives for heterosexual men that should be used in combination with other measures.
2. There is no need or equipoise to conduct a U.S. trial of MC for HIV prevention among men who have sex with women.
3. There is not enough evidence to make a recommendation for MC for MSM to prevent HIV infection and there may be equipoise to conduct an efficacy trial for this population.
4. For newborns, medical benefits outweigh risks and the benefits and risks should be explained to parents.

Role of prepuce in HIV infection

The protective effect of circumcision against HIV infection has been attributed to several factors. Microabrasions during intercourse may provide easier access to the virus and the likelihood of such coital trauma is higher in those uncircumcised.⁵⁸ The inner foreskin mucosa has a higher density of Langerhans cells for which HIV-1 demonstrates specific tropism in in vitro studies.⁵⁹ Circumcision removes a majority of foreskin Langerhan cells. Pro-inflammatory anaerobes, which are supported by the anoxic microenvironment of the intact foreskin, also contribute to viral survival.⁶⁰ The lack of keratinization, especially of the inner foreskin, compared to a circumcised penis may also aid HIV infection.⁶¹ In a sub-analysis in the control group of the Ugandan RCT, foreskin surface area was predictive of the risk of HIV acquisition.⁶² Men with the largest foreskin surface area had a two times higher incidence rate compared to those with the lowest quartile surface area.

In addition, higher rates of STI, such as herpes simplex virus-2 (HSV-2) and genital ulcer disease (GUD), increases susceptibility to HIV infection and circumcision may decrease HIV risk by these intermediate factors, but this relationship is complex and bidirectional.^{63–66}

Female to male HIV transmission

Observational studies

Following the first observational study in 1986 suggesting a decreased risk of HIV in circumcised men, several studies and a meta-analysis of 15 observational studies conducted

in 2000 people, supported a protective effect of circumcision against HIV infection (adjusted risk ratio [RR] 0.42; 95% CI 0.34–0.54%).^{67,68} More recently, Warner et al conducted a cross-sectional study among heterosexual African American men and analyzed the protective effects of MC in a group of men with known HIV exposure.⁶⁹ Circumcision was associated with a 51% reduction in HIV prevalence among the 394 visits by men who were exposed to HIV-positive female partners (adjusted PRR 0.49; 95% CI 0.26–0.93). In contrast, when the HIV exposure was unknown, circumcision status was not protective (adjusted PRR 1.00; 95% CI 0.86–1.15). A global epidemiological study in 118 countries concluded that circumcision was associated with a lower HIV rates, independent of religion.⁷⁰ However, for non-sub-Saharan countries with a primarily homosexual or IV drug use modality of HIV transmission, circumcision status was not associated with a lower risk of HIV infection.

Meta-analysis of HIV RCTs

A Cochrane review of the three trials (Table 1) concluded that medical circumcision reduces the acquisition of HIV by heterosexual men from 38–66% over two years, with an incidence risk ratio of 0.5 (95% CI 0.34–0.72) at one year and 0.46 (95% CI 0.34–0.62) at two years' followup in the circumcised group compared to the non-circumcised group.⁷¹ There was no evidence of heterogeneity (I^2 0%). With a low assumed control risk of 1% (10 per 1000 population over a two-year period), the NNT based on this meta-analysis was 186. The combined per protocol analysis showed a stronger protective effect (incident rate ratio [IRR] 0.34; 95% CI 0.24–0.47).

The possibility of behavioural disinhibition, leading to unsafe sexual practices, can potentially offset the protective effect of circumcision and this effect may be dependent on the timing of circumcision.⁷² The South African trial evaluated behavioural aspects post-circumcision and noted a higher mean number of sexual contacts in the circumcised group compared to the uncircumcised group.⁵⁵ The Kenyan trial also showed a statistically significant difference in the circumcised vs. uncircumcised group with respect to unprotected sexual intercourse and consistent condom usage.⁵³ The Ugandan RCT did not find this difference even on long-term followup.^{54,73} In a comprehensive analysis of 1309 men enrolled in the Kenyan RCT, Mattson et al evaluated risk compensation using a self-validated 18-item risk propensity score and acquisition of other STIs as a marker of risk behaviour.⁷⁴ Men in this study were not clearly informed that MC reduced HIV risk and both groups received intensive counselling. STI risk was higher in the circumcised group at baseline and incident STI was higher at six-month followup. There was no difference in the risk scores at six- and 12-month followup in the two groups.

Male to male transmission

According to CDC estimates in 2006, of 56 300 new HIV infections in the U.S., 53% were in MSM males.⁷⁵ Heterosexual transmission was responsible in 31% of cases. Canadian estimates are similar, with more than 50% of new infections occurring in MSM males.

Evidence of an association between circumcision status and HIV infection in MSM males is limited to observational studies, not usually stratified by receptive and insertive roles. It is believed that men who practice an insertive role during anal intercourse would likely benefit from MC, while those who perform a receptive role have little or no protection. Most MSM are likely to be infected through a receptive rather than an insertive role, which would further dilute the effects of MC in this population. In a Cochrane review by Wiysonge et al, 21 observational studies (six cohort, 14 cross-sectional, one case-control; 71 693 participants) were analyzed to assess the impact of circumcision for preventing HIV infection in MSM males; the risk for HIV acquisition was not associated with MC (OR 0.86; 95% CI 0.7–1.06).⁷⁶ In a subgroup analysis of men reporting an insertive role, MC was found to be protective (3465 participants; OR 0.27; 95% CI 0.17–0.44). The overall quality of the studies included was poor, with high risk of attrition and detection bias. A previous meta-analysis by Millett et al in 2008 had also reported a similar non-significant RR of 0.86 among all MSM men and a trend towards lower HIV risk in MSM men who practiced an insertive role (RR 0.70; 95% CI 0.2–2.2).⁷⁷

Londish et al created a mathematical model incorporating circumcision and seropositioning in the MSM population to predict the reduction in HIV prevalence and incidence.⁷⁸ The authors predicted that in a developed country with 10% HIV prevalence, with universal circumcision, it would take 20 years to reduce HIV incidence by 5% compared to pre-intervention levels and prevalence to 9.6%.

Male to female transmission

MC can potentially decrease female partner HIV infection by a direct effect, or indirectly (over 10–20 years) at the population level by reducing the overall male prevalence.

The Ugandan RCT enrolled 92 couples in the circumcised group and 67 in the control group to study the direct effects of MC in HIV-positive men on female partner HIV status over a 24-month followup.⁷⁹ This trial was underpowered and terminated early since recruitment was futile. No evidence of protection was identified (adjusted HR 1.49; 95% CI 0.62–3.57). A meta-analysis of seven longitudinal, population-based studies, including the Ugandan trial, did not show any protective effect on female partners of circumcised males (RR 0.80; 95% CI 0.54–1.19).⁸⁰ There was evidence of between-study heterogeneity, with studies showing reverse

Table 1. The three randomized trials on circumcision and HIV prevention

Study (location, time period)	Allocation concealment	Blinding	Inclusion criteria	n	Followup	Lost to followup	AEs	Outcomes	ITT RR (95% CI)	As treated RR (95% CI)	Covariate adjusted RR
Auvert et al (South Africa, 2002–2005)	Partial	Study personnel	Male 18–24, wishing to be circumcised	3274	Stopped at 63% of total anticipated person-years	251 (8%), 30% in circumcised and 33% uncircumcised	3.8%	Intervention: 20/1546 Control: 49/1582	0.42 (0.25–0.70)	0.24 (0.14–0.44)	0.39 (0.23–0.66)
Bailey et al (Kenya, 2002–2006)	Unclear	HIV testers, nurses counselling and doing questionnaires partially blinded	HIV-negative, 18–24 years	2784	Stopped at 87% followup	86% (1501) completed 24-month followup, overall 1283/2784 (46%) did not complete trial	1.7%	Intervention: 22/1388 Control: 47/1392	0.47 (0.28–0.78)	0.45 (0.27–0.76)	0.44–0.47
Gray et al (Uganda 2002–2006)	Partial	None specified	HIV-negative, 15–49 years	4996	Stopped at 72% person-time accrual	22% at 24 months followup	8%	Intervention: 22/2387 Control: 45/2430	0.49 (0.28–0.84)	0.45 (0.25–0.78)	0.49 (0.29–0.81)

AE: adverse events; CI: confidence interval; ITT: intention-to-treat; RR: relative risk

effects. In a prospective cohort study based on an RCT conducted to assess HSV-2 suppression impact on HIV transmission conducted in Africa, 1096 serodiscordant couples were followed for an 18-month period.⁸¹ Circumcision was associated with a 40% lower risk of HIV transmission to female partners, although the results were not statistically significant (HR 0.62; 95% CI 0.35–1.10).

Conclusions on the benefit of neonatal circumcision in preventing HIV

There are several factors that have to be considered when adopting findings from recent evidence as a basis for recommendation of neonatal circumcision in more developed countries.^{82–87} The large sample size, magnitude of the effect, consistent results across the three trials and meta-analyses, and sound statistical methods to address confounding factors are the primary strengths of these well-conducted trials, which upholds the internal validity of the results to a large extent barring some concerns. These concerns include early trial stoppage, inadequate allocation concealment and random sequence generation, and risk of attrition bias.^{71,82} The primary issue in interpreting these results relates to external validity of these trials in the Canadian setting.

1. The MC rate in the three African trial sites ranged from 10–20% and HIV prevalence in these countries was from 7–25% (incidence 1.7–1.8% per year). In contrast, in Canada the MC rate is around 35% and HIV prevalence is much lower, and this would lead to a substantially higher NNT.
2. A difference in the modes of HIV transmission and baseline neonatal circumcision rates affects the available susceptible population, which will be protected by MC. Only a small proportion of HIV transmission is due heterosexual activity and MSM is a group not protected by MC.
3. Variations in sexual practices and behaviour (including condom usage) and differences in STI prevalence will also alter the protective effects of MC.
4. Access to healthcare and earlier detection and treatment for HIV-infected males and HPV vaccination programs may also modify the observed protective effects.
5. Ethical considerations of parental consent and racial/ethnic acceptability further complicate the issue when implementing universal circumcision programs.
6. Cost benefit analysis compared to alternative preventive strategies should be considered and studied in a Canadian context to allow generation of a clear recommendation.⁸⁷
7. The trials were all conducted in sexually active adult men from HIV-endemic areas in Africa, who were motivated and interested in a free circumcision. They also received counselling as part of the trial. This will result in overestimation of the protective effect, different from a more real-world setting.
8. In addition, the long-term effectiveness beyond two years' followup is currently only published for the Ugandan trial.⁷³

Recommendations (MC and HIV protection):

1. **Female to male transmission: There is compelling evidence that MC reduces the risk of HIV transmission from female partners to male (Level 1 a, Grade A). The magnitude of the effect is debat-**

able and cannot be extrapolated to Canada from the African RCTs.

2. **Male to male transmission: Based on current evidence, MC does not provide protection for men who have sex with men (Level 2a).**
3. **Women partners: Based on current evidence, MC is not protective for female partners (Level 2a–b).**
4. **Universal infant circumcision cannot be recommended to prevent HIV infection based on current evidence (Grade B).**

2. Circumcision and HPV prevention

HPV is the commonest STI worldwide and of the more than 100 types, about 40 can infect the ano-genital area. In the absence of vaccination, up to 75% of Canadians would have at least one lifetime HPV infection (www.phac-aspc.gc.ca). High-risk oncogenic types, like 16 and 18, are implicated in cervical, penile, vulval, vaginal, anal, and some oropharyngeal cancers, while low-risk non-oncogenic subtypes, like 6 and 11, cause genital warts. The effect of circumcision on HPV is difficult to interpret, as HPV infection can be transient, affect multiple genital areas outside the foreskin, include several high-risk and non-high-risk types, and is significantly associated with other behavioural confounders. In addition, HPV prevalence, incidence, clearance, and viral load are all potential outcomes that can be studied and have differing health implications.

HPV in men

Level 1–2 evidence

Auvert et al showed a reduction in the prevalence of urethral high-risk HPV infection following male circumcision, with a prevalence rate ratio of 0.68 (95% CI 0.52–0.89; $p=0.002$) in circumcised men as compared to uncircumcised men.⁸⁸ Significantly, the prevalence differences between the two groups were not significant for HPV type 16, but were for HPV type 18.

Six secondary trials analyzed HIV-positive and -negative men enrolled in the Ugandan HIV trial with regards to HPV prevalence, acquisition/incidence, clearance, and viral load.^{89–94} The first trial investigated the prevalence of HPV in a subgroup of participants and only included samples from the glans and coronal sulcus, a factor for possible bias due to differential infection sites in circumcised males.^{89,95} The adjusted risk ratio for prevalence of high risk HPV at two years' followup in the circumcised group was 0.65 (95% CI 0.46–0.90; $p=0.0009$) and 0.66 for low-risk HPV genotypes (95% CI 0.49–0.91; $p=0.01$). In a subsequent report, the same group showed that the one-year penile shaft HPV prevalence after MC was not statistically significantly

lower in the circumcised group (adjusted PRR 0.66; 95% CI 0.39–1.12; $p=0.12$).⁹⁰

The third trial evaluated HPV acquisition and clearance in HIV-negative men using glans and coronal sulcus samples.⁹¹ The incidence rate of HR HPV infection was statistically significant at one year (IRR 0.61; 95% CI 0.44–0.85) in the uncircumcised group, but at two years' followup, the effect was not statistically significant (IRR 0.64; 95% CI 0.38–1.07). The incidence of type-specific HR HPV was statistically significant for only the 18 and 33 genotypes and not 16. The acquisition of new multiple infections was lower in the circumcised group (IRR 0.45; 95% CI 0.28–0.73) compared to the non-circumcised group. Clearance rates per 100 person years were statistically significant only for types 39, 51, and 58 and overall clearance rates were higher in the circumcised group (RR 1.36; 95% CI 1.13–1.63). The fourth study in the Ugandan trial evaluated HR HPV prevalence, acquisition/incidence and clearance in HIV-positive married men with or without a circumcision.⁹² A random sample comprising 22% of those enrolled was tested at enrollment and at 24 months' followup. MC provided partial protection, with the circumcised group (55% positive) showing lower HR HPV prevalence at 24 months (PRR 0.77; 95% CI 0.62–0.97). The incidence rate for one or more new infections after adjustment was not statistically significant between the circumcised and non-circumcised group (IRR 0.74; 95% CI 0.54–1.01); however the proportion of men acquiring multiple new HR HPV infections was lower in the circumcised group (IRR 0.40; 95% CI 0.19–0.84). The clearance rate of HPV infections was not significantly different between the two arms. In a more recent analysis of 999 men (HIV-positive and -negative) from the Ugandan trial, Tobian et al showed an increased HPV clearance in HIV-negative circumcised men (adjusted RR 1.48; 95% CI 0.55–0.89) and lower incidence of HR HPV acquisition in HIV-positive men (IRR 0.70; 95% CI 1.67–2.44).⁹³ The final study evaluated HPV viral load in circumcised and uncircumcised HPV-infected men at 24 months.⁹⁴ MC decreased HPV viral load in circumcised compared to uncircumcised men for new infections acquired after enrollment, but the results were statistically significant for only serotype 16 ($p=0.001$).

Two meta-analyses were conducted evaluating the effect of MC on HPV.^{96,97} Albero et al in 2012 analyzed data from 14 observational studies and two RCTs conducted between 1971 and 2010.⁹⁷ Accepting heterogeneity in MC reporting status, sites sampled and methods of detection, in the 1784 participants analyzed with data from the two RCTs, the authors detected a strong inverse association between circumcision and high-risk HPV prevalence, with an OR of 0.67 (95% CI 0.54–0.82). The 14 prevalence studies showed a similar pooled result of overall HPV prevalence (OR 0.57; 95% CI 0.42–0.77). HPV prevalence remained lower in circumcised men, even pooling studies where the penile shaft

or scrotum was sampled. There was no association found between circumcision and new genital HPV infections or clearance. A previous meta-analysis conducted by Larke et al until 2010 also showed similar prevalence results (OR 0.57; 95% CI 0.45–0.71).⁹⁶ This prevalence difference diminished at sites away from the glans and urethra. There was weak evidence that circumcision was associated with decreased HPV incidence (RR 0.75; 95% CI 0.57–0.99) or clearance (RR 1.33; 95% CI 0.89–1.98).

Level 2c–4 evidence

The evidence in regards to effectiveness of circumcision in prevention of HPV transmission is contradictory when assessing observational and ecological studies. Dickson et al followed 450 children from birth in a cohort study conducted in New Zealand up to 32 years of age, with circumcision status reported by mothers at three years of age.⁹⁸ Sexual behaviour was recorded at 21, 26, and 32 years, along with assessment of socioeconomic and moral-religious emphasis of family. Seropositivity for HPV 16 or 18 at 32 years was lower in the uncircumcised group (adjusted OR 1.4; 95% CI 0.85–2.2), but more associated with lifetime number of partners and moral-religious emphasis of the family of origin. Albero et al reported on 3969 participants from three countries followed over four years and used coronal sulcus/glans, penile shaft, and scrotal swabs to compare HPV prevalence in circumcised and non-circumcised males.⁹⁹ A multivariable analysis adjusting for race, marital status, lifetime female sexual partners, female sexual partners in 3–6 months, and male sexual partners in the past three months did not find any association between MC and oncogenic HPV (PR 0.95; 95% CI 0.87–1.03). MC was significantly associated with a decreased risk of non-oncogenic HPV infection. Circumcision was associated with a significantly lower HPV 16 seroincidence in MSM males reporting an insertive role during sex (adjusted HR 0.47; 95% CI 0.28–0.98; $p=0.043$).¹⁰⁰

HPV in female partners

Two trials conducted on HIV-positive and -negative men in Uganda analyzed transmission of HPV to female partners.^{101,102} The implications of reduced HPV infection in female partner cervical cancer rates are not clearly discernible since there are several other confounding risk factors.

In the first trial on HIV-negative men and their partners, the two-year prevalence of HR HPV infection in partners was 28% in the circumcised group and 39% in the uncircumcised group (PRR 0.72; 95% CI 0.60–0.85).¹⁰¹ The incidence of any HR HPV infection from 0–2 years was also lower in the circumcised group female partners (IRR 0.77; 95% CI 0.63–0.93). In terms of specific HR genotypes, the results

were specifically not statistically significant for HR HPV 16 or 18. The clearance rate for all genotypes was 66% in the circumcised group partners as opposed to 59% in the uncircumcised partners ($p=0.014$). This clearance rate was reversed for HPV 16, with the uncircumcised group having a 74% clearance rate vs. 52% in the circumcised group (RR 0.70; 95% CI 0.54–0.92). In a second trial on female partner HPV infection, Tobian et al studied the effect of MC in HIV infected men.¹⁰² Circumcision status in male partners was not associated with lower HR HPV prevalence in female partners (PRR 1.07; 95% CI 0.86–1.32; $p=0.64$) or lower HR HPV incidence over two years (IRR 1.05; 95% CI 0.77–1.43) or clearance rates (RR 0.96; 95% CI 0.83–1.11).

Alternative strategies in HPV prevention

The benefit of MC on HPV infection dynamics has to be analyzed in the presence of complimentary HPV vaccination, protection offered by routine condom usage, and other safe sexual practices. HPV vaccination is currently available and recommended for males (HPV 4 vaccine, 9–26 years of age), females (HPV 2 or 4 vaccine, 9–13 years and 14–26 years of age) and MSM (HPV 4 vaccine, >9 years of age) with good evidence (*Level 1, Grade A*) of its effectiveness and safety (National Advisory Committee on Immunization, HPV guideline 2007). A prospective cohort study showed that correct and consistent condom usage also decreased risk of HPV transmission by 70% in young, newly sexually active women.¹⁰³

Recommendations (MC and HPV infections)

1. **HPV prevalence in men: Current evidence suggests a modest decrease in HPV prevalence in the glans and coronal sulcus up to two years following MC (*Level 1b*). The protective effect is partial, does not cover all high-risk types and is weaker further away from the glans and coronal sulcus. It is not clear whether this effect will persist into adulthood following neonatal circumcision.**
2. **HPV clearance in men: There is no evidence (except a single RCT on HIV-negative men) that MC increases HPV clearance (*Level 1b–2b*). If it did increase clearance, this may also inflate the impact of the prevalence benefits mentioned.**
3. **HPV incidence or acquisition in men: There is no convincing evidence to suggest that MC decreases HPV acquisition or incident infections in HIV-positive or -negative men (*Level 1b–2b*).**
4. **HPV in female partners: MC lowers prevalence and incidence in partners of HIV-negative men and improves clearance rates (*Level 1b–2b*).**
5. **As a public health intervention, it is likely that the effect of HPV vaccination and behavioural modification will be more effective than performing universal neonatal circumcisions on all males (*Grade B*).**

3. Circumcision and non-ulcerative STI prevention

The most common non-ulcerative STIs are gonorrhoea, Chlamydia and Trichomonas infections. Chlamydia is the most commonly diagnosed bacterial STI, with approximately 65 000 cases reported in Canada in 2006. These STIs are initiated by bacterial binding to a variety of host receptors and unlike HIV, a biological explanation of how circumcision can be protective against these infections is lacking.

Two RCTs have addressed the role of MC in these infections. In the Kenyan study, there was no association between circumcision status and non-ulcerative STIs, but condom usage was protective (HR 0.64; 95% CI 0.50–0.82).¹⁰⁴ The Orange Farm study showed lower *Trichomonas vaginalis* (*T. vaginalis*) infection in men only in an as-treated analysis (adjusted OR 0.47; 95% CI 0.25–0.92).¹⁰⁵ A meta-analysis of 30 observational studies failed to identify a statistically significant association between non-ulcerative STIs and MC.¹⁰⁶

In a prospective, multicentre, U.S. study involving 2021 men, there was evidence for a statistically non-significant increased prevalence and incidence of gonorrhoeal infection in non-circumcised men (OR 1.3; 95% CI 0.9–1.7 and 1.6; 95% CI 1.0–2.6, respectively), but no difference with respect to Chlamydia infection.¹⁰⁷ In another prospective study, uncircumcised male partners had a higher risk of *T. vaginalis* infection compared to circumcised partners of *T. vaginalis*-infected women (OR 1.8; 95% CI 1.1–3.2).¹⁰⁸

Mycoplasma genitalium can cause urethritis, cervicitis, and pelvic inflammatory disease. Multivariate analysis of data collected in the Kenyan HIV trial showed a higher likelihood of *M. genitalium* infection in non-circumcised men (adjusted OR 0.54; 95% CI 0.29–0.99).¹⁰⁹ Washing the penis within the first hour after sex had a protective effect. In a prospective cohort study, circumcision did not seem to have any protective effect on female partners with regards to Chlamydial, gonococcal, and Trichomonal infections.¹¹⁰

Recommendations (MC and non-ulcerative STIs):

1. **Currently, there is no significant evidence to support the protective role of MC in the acquisition of non-HPV, non-ulcerative STIs (Level 2a–b, Grade B).**

4. Circumcision and prevention of GUD and ulcerative STIs

HSV-1 and -2, *T. pallidum* (syphilis), *H. ducreyi* (chancroid) and *K. granulomatis* (Donovanosis) are the common causes of GUD, with HSV infections accounting for 70–80% of the infections leading to a genital ulcer (Public Health Agency of Canada). The true incidence of HSV-1 and -2 is unknown, but these infections are very common, with estimates based on serological tests suggesting at least 20% prevalence in Canada (Public Health Agency of Canada). Chancroid is extremely rare in Canada and acquisition is primarily limited to endemic areas. Previously rare in Canada, the incidence

of syphilis has shown an increase, especially in MSM males and sex workers.

Women and men with GUD and HSV-2 have a higher risk of acquiring or transmitting HIV and conversely HIV infection increases the risk of GUD.^{111,112} Observational studies suggest that prevalent HSV-2 infection leads to a 2–3-fold increase in the risk of HIV acquisition, and this risk increases up to 7-fold with incident HSV-2 infection.^{113,114} In a HIV vaccine trial (Step study) conducted in MSM men, HSV-2 infection was an important risk factor for HIV acquisition among vaccine and placebo recipients (HR 2.2; 95% CI 1.4–3.5).¹¹⁵ It is likely that the correlation between these two infections is due to high-risk sexual behavior, making it unclear whether HSV-2 acquisition is an important direct cofactor for HIV infection.^{116–118}

HSV

Condom usage has a limited role in preventing HSV transmission, with a 50% protective rate for male to female transmission. Unlike HIV, HSV transmission is less dependent upon the presence of foreskin mucosa.

Level 1–2 evidence

In the Ugandan RCT, which included HIV-positive men, Tobian et al showed a lower risk of HSV-2 seroconversion in the circumcised group over two-year followup (adjusted IRR 0.70; 95% CI 0.55–0.91).¹¹⁸ Consistent condom usage had a slightly higher protective effect (adjusted IRR 0.56; 95% CI 0.36–0.89). In a second analysis of the Ugandan RCT with HIV-negative men, the partial protective effect of MC against HSV-2 seroconversion was similar.⁸⁹ Multivariate analysis of South African RCT data did not show a protective effect for MC against HSV-2 seroincidence (IRR 0.68; 95% CI 0.38–1.22), but the effect was reversed in an as-treated analysis, presumably due to a 8.2% crossover rate (IRR 0.45; 95% CI 0.24–0.82).⁶³

Mehta et al conducted a RCT in Kenya to assess the protective effect of circumcision against HIV, HSV-2 and GUD.⁶⁵ HSV-2 incidence did not differ by circumcision status (RR 0.94; 95% CI 0.7–1.25), but HSV-2 incident infection tripled the risk of HIV acquisition (RR 3.44; 95% CI 1.52–7.80). Moreover, non-HSV-GUD risk was reduced by 50% in those circumcised (RR 0.52; 95% CI 0.37–0.73). In a multivariable model, the presence of GUD was associated with a seven times greater risk of HIV seroconversion, suggesting that the protective effect of circumcision may be partially mediated by reducing the risk of GUD. More than 50% of HIV seroconversions were preceded by HSV-2 or GUD.

Level 2–4 evidence

In a meta-analysis of observational studies, MC was not associated with a decreased risk of HSV-2 seropositivity (RR 0.88;

95% CI 0.77–1.01) and this effect was less protective when restricting the analysis to studies using genital examination rather than self-reported circumcision status (RR 0.97; 95% CI 0.80–1.17).¹⁰⁶ There was a protective effect of MC on syphilis seropositivity (RR 0.67; 95% CI 0.54–0.83), although there was significant heterogeneity among the studies and two of the largest studies included in this analysis showed the least protective effects. A definitive conclusion could not be reached for the risk of chancroid, with the adjusted RR varying from 0.13–1.11.

Xu et al estimated the prevalence of circumcision in the U.S. and examined the association between MC and HSV-2 infection using the National Health and Nutrition Examination survey conducted on 6174 men.¹¹⁹ HSV-2 infection was associated with age, race, and sexual behaviours, but not with circumcision status (OR 1.1; 95% CI 0.8–1.5).

Male to male HSV-2 transmission

In an observational study on 3828 men, Jameson et al found that, even in men reporting primarily an insertive role, MC was not protective against HSV-2 (adjusted OR 0.66; 95% CI 0.27–1.63).¹²⁰ Barnabas et al conducted a cross-sectional study on MSM males and found that MC was associated with a borderline protective effect against HSV-2 infection (OR 0.7; 95% CI 0.5–1.0).¹¹⁴ In a Cochrane review of MSM males, circumcision did not have a protective role in preventing syphilis (OR 0.96; 95% CI 0.82–1.13) or HSV-2 infections (OR 0.86; 95% CI 0.62–1.2).⁷⁷ In developed countries, the results of MC can be contradictory, with opposite direction of the effect shown in two longitudinal cohort studies from New Zealand.^{121,122}

Non-ulcerative and ulcerative STIs in female partners

Gray et al conducted a sub-trial in the Ugandan HIV RCT on 1563 HIV-negative women married to men randomized to circumcised and non-circumcised groups followed for a year.¹²³ Adjusted analyses suggested a 22% circumcision efficacy for GUD (adjusted PRR 0.78; 95% CI 0.61–0.99), a 45% efficacy for trichomonas (adjusted PRR 0.55; 95% CI 0.34–0.89), and a weak 18% efficacy for bacterial vaginosis (BV) (adjusted PRR 0.82; 95% CI 0.74–0.91). In the Ugandan RCT on HIV-negative men, MC did not reduce the risk of HSV-2 acquisition in women partners of both HSV-2-negative and -positive male partners.¹²⁴ In addition, a 2008 prospective study of almost 6000 HIV-negative women showed no difference in female acquisition of Chlamydia (adjusted HR 1.25; 95% CI 0.96–1.63), gonorrhea (adjusted HR 0.99; 95% CI 0.80–1.36) or *T. vaginalis* (adjusted HR 1.05; 95% CI 0.80–1.36) according to circumcision status.¹¹⁰

Recommendations (MC and ulcerative STIs):

1. Currently, there is no significant evidence to support the protective role of universal neonatal cir-

cumcision for males and females in the acquisition of ulcerative STIs (Level 2–4, Grade C).

2. There is weak evidence of decreased seroconversion for HSV-2 following MC in adult men in Africa (Level 2a–b).

Circumcision and risk of penile cancer

Epidemiology

Penile cancer is a rare disease with age-standardized incidence rates of 0.3–1.0 per 100 000 men in Europe and North America, accounting for 0.4–0.6% of all malignancies.¹²⁵ The potential risk factors for penile cancer include phimosis and balanitis, smoking, HPV infections, penile oral sex, Lichen sclerosis, premalignant conditions like Bowen's disease and erythroplasia of Queyrat, priapism, urethral stricture, and psoralen and ultraviolet A (PUVA) therapy.^{126–128} Therefore, MC for preventing penile cancer is possibly one of the several preventive interventions, which still does not account for a host of risk factors.^{127–129} The International Consultation on Urologic Disease 2009 consensus publication on penile cancer prevention advocated smoking cessation as a Grade C recommendation, while male HPV vaccination and universal circumcision were not recommended.¹²⁸

Biological plausibility

Penile cancer is mediated through two mechanisms: HR HPV infections and a subset through non-HPV-mediated mechanisms related to phimosis and Lichen sclerosis. Oncogenic HPV (mainly type 16 and 18) prevalence is noted in 40–45% of penile cancers in several systematic reviews.^{130–134} In a Belgian study, HPV DNA was identified in 61% of invasive penile cancer samples, with the commonest serotypes being HPV 16, 11, 56, and 18.¹³⁴ Importantly, only 13% of the cases in this study were infected with HPV types present in HPV vaccines.

Level 2 evidence

In a meta-analysis of eight studies (seven case control, one cross-sectional study), Larke et al showed that circumcision at <18 years of age was protective against invasive penile cancer with an OR of 0.33 (95% CI 0.13–0.83).¹³³ In those circumcised as adults, the risk of invasive carcinoma was increased (OR 2.71; 95% CI 0.93–7.94), presumably because surgery was performed for conditions predisposing to penile cancer.

Level 3 evidence

In a matched case control study, Tsen et al showed that phimosis was a strong risk factor for invasive penile cancer (adjusted OR 16; 95% CI 4.5–57).¹³⁵ The protective effect of neonatal circumcision was not statistically significant when the analysis was restricted to those who did not have a history of phimosis (OR 0.79; 95% CI 0.29–2.6) and smoking was a clear identified risk factor (OR 5.9 for >20 cigarettes/day). In another population-based case control study from Denmark, Madsen et al found that penile cancer was positively associated with measures of high and early sexual activity, genital warts, unprotected sex, and penile oral sex.¹³⁶ Phimosis (OR 4.9; 95% CI 1.85–13.0), but not childhood circumcision ($p=0.33$) was also found to have a strong association on multivariate analyses. The authors concluded that an unretractable foreskin with HR HPV infection might constitute the single most important risk factor for penile cancer. In a population-based case control study by Daling et al, 137 men with penile cancer were compared with 671 controls.¹³⁷ Lack of childhood circumcision (OR 2.3; 95% CI 1.3–4.1), phimosis (OR 7.4; 95% CI 3.7–15.0), and cigarette smoking (OR 4.5; 95% CI 2.0–10.1) were identified as risk factors for invasive penile cancer, but after excluding patients with phimosis, the analysis did not show a protective effect of childhood circumcision.

Level 2c evidence

Indirect evidence of the role of MC in preventing penile cancer can be investigated by ecological studies in countries with low circumcision rates. Denmark, with 2% circumcision prevalence, showed decreasing and lower rates of penile cancer than in the U.S.¹³⁸ However, in a more recent study from Denmark, there was an increase in the incidence of penile cancer from 1 to 1.3 per 100 000 men-years between 1978 and 2008.¹³⁹ In the U.S., despite a decrease in circumcision rates, a 1.2% average annual incidence rate decrease has been noted between 1973 and 2003.¹⁴⁰

Recommendations (MC and penile cancer):

1. **Circumcision decreases the risk of penile cancer (Level 2–3).**
2. **However, given the low incidence of invasive penile cancer, the partial protective effect of MC, and the availability of other preventive strategies, such as HPV vaccination, condom use, and smoking cessation programs, it is difficult to justify universal neonatal circumcision as a preventive strategy for preventing penile cancer (Grade B).**
3. **Recognition and treatment of phimosis during regular health visits is recommended to decrease the risk of penile cancer (Level 5, Grade D). A genitourinary exam during puberty is recommended to ensure pre-**

putial retractability and hygiene, rule out phimosis, and counsel regarding HPV vaccination and safe sexual practices, as well as to offer the possibility of circumcision as a preventive measure against STIs while specifying the drawbacks and efficacy of other preventive measures (Grade D).

Circumcision and risk of prostate cancer

A meta-analysis of case control studies by Taylor et al found an increased RR of prostate cancer in men with a history of STIs.¹⁴¹ A recent case control study explored the association between circumcision and prostate cancer.¹⁴² In a multivariable analysis, controlling for age, family history, race, history of STIs, number of partners, and history of prostatitis, the authors did not find an overall association with circumcision (OR 0.87; 95% CI 0.74–1.02). A previous case control study from the U.K., looking primarily at dietary and sexual history, found a borderline association on univariate analysis between circumcision and prostate cancer risk (OR 0.62; 95% CI 0.39–0.98).¹⁴³

Conclusion (MC and prostate cancer): There is no convincing evidence on the protective effect of MC against prostate cancer (Level 3–4, Grade B).

Role of the foreskin in sensation and sexual function

There is ongoing controversy regarding the impact of circumcision on penile sensitivity and sexual satisfaction. It is obvious that the foreskin has sensory nerves, which are lost following a circumcision. The primary question is whether this presumed loss of sensation or a possible decrease in glans sensitivity impacts sexual satisfaction in a measurable and consistent way after accounting for several confounders. The timing of circumcision (adult vs. neonatal) may also impact this effect. The problem is amplified by the lack of a single objective measure of sensitivity (sensation varying by type and site assessed, ejaculation latency, arousal). In addition, this effect of MC has to be studied both from the men and their male and female partners' perspective. It is hard to extrapolate results of adult MC studies on sexual function and sensation to neonatal circumcision.

Adult circumcision

Level 1–2 evidence

A recent meta-analysis included 10 studies with significant heterogeneity and poor methodological quality to assess the impact of MC on sexual function.¹⁴⁴ There were no significant differences in sexual desire, dyspareunia, premature ejaculation, ejaculation latency time, or erectile dysfunctions between circumcised and uncircumcised men. A secondary

analysis of the Ugandan RCT showed no long-term differences in 4456 men randomized to immediate and delayed circumcision arms who were assessed at six, 12, and 24 months for sexual desire, satisfaction, and erectile dysfunction.¹⁴⁵ Although self-reported, there were some significant differences in penetration and pain on intercourse noted at six months favouring uncircumcised men, but this difference normalized over the followup period. The trial showed that there was a higher improvement in sexual satisfaction in the uncircumcised group compared to the circumcised group. In comparison, another trial from Kenya noted that circumcised men reported increased penile sensitivity and enhanced ease of reaching orgasm with no sexual dysfunction as compared to uncircumcised controls.¹⁴⁶ Sexual dysfunction decreased significantly in both the circumcised and uncircumcised men during followup. The two trials reported very different baseline sexual dysfunction and, in both, the uncircumcised group reported improvement in sexual satisfaction over time. The Kenyan RCT showed a reduced risk of any self-reported coital injury in those circumcised compared to uncircumcised men (OR 0.61; 95% CI 0.54–0.68).¹⁴⁷ There was a significant decrease in reported penile injuries over followup even in the control uncircumcised group (decrease from 64% to 43%).

Level 3–4 evidence

In a survey conducted in Denmark, there was no difference in the circumcised and uncircumcised group in current sexual activity, but circumcised men were more likely to report orgasm difficulties (adjusted OR 3.26; 95% CI 1.05–4.16).¹⁴⁸ In addition, women partners with circumcised spouses also noted orgasm difficulties (adjusted OR 2.66; 95% CI 1.07–6.66) and dyspareunia (adjusted OR 8.45; 95% CI 3.01–23.74). Another cross-sectional study conducted in Belgium on 1369 men (1059 uncircumcised, 310 circumcised) used a self-reported online version of the Self-assessment of Genital Anatomy, and Sexual Function, Male questionnaire (SAGASF-M) to measure four dimensions of sexual function.¹⁴⁹ Overall, for the penis as a whole, the two groups differed in sexual pleasure ($p=0.044$) and discomfort/pain ($p=0.018$), both favouring the uncircumcised group. The study concluded that circumcision led to a decrease in glans sensitivity and overall penile sensitivity. In another detailed study of fine-touch pressure thresholds in circumcised and uncircumcised men, Sorrells et al noted that the glans of uncircumcised men had significantly lower pressure thresholds compared to circumcised men when adjusted for age, type of underwear, and ethnicity.¹⁵⁰ Studies before and after circumcision in the adult population generally do not show any differences in sexual activity and function, although these studies are often hard to interpret because of the small sample size, lack of validated instruments to measure sexual function, self-reported outcomes, short fol-

lowup times after circumcision, and presence of medical indications for circumcision.^{151–154}

Intravaginal ejaculation latency time (IELT) was measured in two multinational studies using a stopwatch and a blinded timer.^{155,156} The significance of IELT as a measure of sexual satisfaction and sensation is debatable, as a high IELT may suggest a low sensation and conversely a low IELT may suggest premature ejaculation and eventual lower sexual satisfaction. Circumcision and condom use did not impact IELT in both studies.

Neonatal circumcision

A single study compared men circumcised in the neonatal period to uncircumcised men with normal and abnormal erectile function and used quantitative somatosensory testing for assessing glans sensation.¹⁵⁷ Circumcised men with or without erectile dysfunction had worse vibration and better pressure thresholds, but these differences disappeared when controlled for age, hypertension, and diabetes.

Sexual function in partners

In an analysis of self-reported sexual experience in women partners of men who participated in the Ugandan RCT before and after circumcision, Kigozi et al showed no changes in 57% and improved sexual satisfaction in 40%.¹⁵⁸ In 25% of the women who reported better sexual satisfaction, the reason given was related to the male seeking more frequent sex. In a study comparing uncircumcised and circumcised homosexual men as part of the HIM (Health in Men) cohort, Mao et al noted no differences in sexual difficulties or type of anal sex practiced.¹⁵⁹ Men circumcised after infancy were more likely to practice receptive anal sex and had a higher incidence of erection difficulties.

Conclusion: There is lack of any convincing evidence that neonatal circumcision will impact sexual function or cause a perceptible change in penile sensation in adulthood (Level 3–4, Grade C).

Medical indications for childhood circumcision

Pathological phimosis

Pathologic phimosis is an uncommon pediatric diagnosis (0.6–1.5% of boys) and is diagnosed by the presence of a whitish, fibrotic preputial ring.¹⁶⁰ This is different from physiological phimosis, where gentle retraction during examination will show “flowering” or pouting of the preputial orifice and lack of the cicatricial ring.^{161–164} Alternative treatments, such as preputioplasty, dorsal slit, or steroid therapy, can be attempted, but depending on the severity of the scar tissue,

circumcision may be the only curative option when true phimosis is diagnosed.¹⁶⁵⁻¹⁶⁶

Genital lichen sclerosis

Genital lichen sclerosis (LS) or balanitis xerotica obliterans is a chronic, inflammatory dermatosis of the prepuce and glans penis, which can potentially involve the meatus and urethra. The etiology is unknown and probably multifactorial, with a possible autoimmune or infective etiology.¹⁶⁷ The disease tends to be progressive and affect older children; those with obesity and previous surgery tend to have more severe disease.¹⁶⁸ LS should be suspected when clinical examination reveals a more impressive (than phimosis) thick white ring-like cicatrix at the distal preputial ring, associated with white discoloration and plaque formation. A history of secondary phimosis in a child with a previously retractile foreskin and failure of topical steroid therapy is also highly suggestive of LS.¹⁶⁸⁻¹⁷⁰ Overall, the incidence of meatal involvement leading to stenosis is low and estimated to be around 2%.^{169,170} Meatal or urethral involvement is more likely with a history of previous surgery and was seen in 27% in the series by Gargollo et al.¹⁶⁸

The incidence of this condition is underestimated and recent evidence suggests that in boys referred with a diagnosis of phimosis, the incidence ranges from 10–40%.¹⁶⁹⁻¹⁷² In a series of 100 boys referred for phimosis, the incidence of LS was 1.8% under six years of age and up to 21% in those older.¹⁷¹ In another prospective 10-year study by Kiss et al, the incidence of histologically confirmed LS was 40%, and 93% of LS patients had a history of secondary phimosis.¹⁷⁰ In another series of consecutive patients from the U.K., Yardley et al noted a 34% incidence of LS in boys who underwent circumcision and an overall 12% prevalence of LS in boys referred to a specialist for foreskin problems.¹⁷³ The pathological diagnosis of LS may not correlate with clinical suspicion, suggesting circumcision specimen should be routinely subjected to histological examination to rule out LS.¹⁷⁴

The use of topical steroids in LS is debatable, with low response rates. It also requires close followup, as disease progression may lead to glans and urethral involvement.^{167,170,175} Circumcision is usually curative, but some children, depending on the degree of involvement, may need a meatoplasty, glans resurfacing, or urethral reconstruction.¹⁶⁸ If the meatus is clearly involved, a meatoplasty is indicated during circumcision; however, if the involvement is doubtful, close followup during the post-operative period and uroflowmetry assessments for up to two years may be indicated to rule out stenosis.¹⁷⁶ In addition, secondary to the Koebner phenomenon, recurrent LS may appear along scar lines of previous surgery; the use of topical steroids is an option in the postoperative period.¹⁶⁷

Recurrent UTIs

Circumcision can be performed as an adjunct or alternative to prophylactic antibiotics in infants with UTI-predisposing urological abnormalities, as described in the section on UTIs.

Contraindications of neonatal circumcision

Neonatal circumcision should be performed on medically stable, term infants without other medical conditions that require ongoing management or increase risk of surgery. Routine neonatal circumcision should not be carried out in children with congenital anomalies of the penis, including hypospadias or epispadias (Figs. 4, 5), penoscrotal webbing (Fig. 6), concealed penis (Fig. 7), and ventral curvature (Fig. 8). In some of these conditions, a circumcision can be performed with appropriate technical modifications, but this requires a pediatric urological consult. Prior circumcision may not compromise distal hypospadias repair in children. In addition, some children with a hypospadias variant termed megameatus-intact prepuce hypospadias have a normal foreskin and a distal hypospadias only uncovered during a circumcision (Fig. 9).¹⁷⁷ Most of the children with this variant or a distal hypospadias can proceed with a circumcision; however, this requires an ability to recognize the severity of the anomaly and, therefore as a general rule, all boys with hypospadias should ideally not have a circumcision prior to a consultation by a pediatric urologist. Children with blood dyscrasias can undergo circumcision, under appropriate treatment and care.¹⁷⁸⁻¹⁷⁹

Anesthesia for neonatal circumcision

It is clear that neonatal circumcision must be performed with adequate anesthesia and analgesia.^{180,181} The adverse physiological and behavioural responses of inadequate pain control in neonates is convincing, can lead to potential complications, and can alter long-term pain responses in the neonate.¹⁸⁰⁻¹⁸² Different methods used for providing anesthesia and/or analgesia during circumcision include general anesthesia, topical anesthetics, penile nerve blocks, oral sucrose-glucose administration, non-nutritive sucking, caudal block, and various combinations of the above. In addition, the timing of neonatal circumcision may impact pain scores, with earlier surgery being beneficial.¹⁸³ Despite a standardized technique, Neonatal/Infant Pain Scores increased significantly beyond eight days of life, and all neonates beyond three weeks of age had a score indicating pain during the procedure.

There are three topical anesthetic options currently available: lidocaine-prilocaine 5% cream (EMLA), tetracaine 4% gel, and liposomal lidocaine 4% cream. When compared to placebo, crying time was shortened and the heart rate



Fig 4. Hypospadias.



Fig 5. Epispadias.

reduced in children who were circumcised under EMLA.¹⁸⁴ The analgesic effect of EMLA takes almost 60 minutes to take effect and alone may not be sufficient to control the pain arising from foreskin removal. Wahlgren et al noted that the depth of penetration is variable with EMLA and increases with application time to approximately 6 mm after 3–4 hours.¹⁸⁵ Reactions to EMLA included erythema and blanching. Increased methaemoglobin levels secondary to oxidation of hemoglobin by prilocaine metabolites was found to be within normal limits in two trials of EMLA.¹⁸⁵ Liposomal lidocaine is available in the U.S. and has a shorter onset of action.

Dorsal penile (DPNB) and ring blocks are effective techniques to manage circumcision-related pain. Based on a RCT that compared these two methods with EMLA for neonatal circumcision, there was no statistical difference in crying time and heart rate between a ring and DPNB, while EMLA was less effective.¹⁸⁶ Two other trials comparing EMLA to DPNB also demonstrated significantly lower behavioural

distress scores and Neonatal Infant Pain Scale scores in the DPNB group.^{187,188} Cyna et al did not find a difference between caudal blocks and DPNB in a meta-analysis of five RCTs comparing these interventions, but highlight the possibility of a motor block and requirement of anaesthesia expertise make a caudal block less preferable in older ambulatory children.¹⁸⁹ Based on the Cochrane review by Brady-Fryer, a DPNB is the most effective intervention for circumcision-related pain with the caveat that the injection is performed appropriately.¹⁸⁰ A ring block has similar efficacy and may be easier and safer to use. EMLA cream and other topical anesthetics are an option when expertise with penile nerve blocks is not available, and ideally should be used in conjunction with a block.

The DPNB is performed by injecting subcutaneously at the 11 and 1 o'clock positions on the dorsum of the penis close to the base of the penis using a 25-gauge needle. It is important to aspirate prior to injection to prevent intracor-



Fig 6. Peno-scrotal webbing.



Fig 7. Concealed penis.



Fig 8. Ventral curvature.

poreal or dorsal vein injection. A wait time of 5–8 minutes is recommended to achieve adequate anesthesia. Warming the lidocaine or injecting slowly can decrease the pain associated with injection. In a series of 3909 DPNBs, the overall complication rate was 0.18%.¹⁹⁰ Long et al studied the sensory distribution of the penile skin and recommend a ventral infiltration just proximal to the ventral foreskin incision to add to the DPNB.¹⁹¹

Although it has been shown that analgesic methods, such as oral administration of sucrose, glucose, or parenteral acetaminophen, were more effective than placebo, it is widely accepted that these methods are not sufficient as sole measures for relieving the pain associated with circumcision.^{180,192} South et al suggested addition of non-nutritive sucking to DPNB based on a RCT, which showed significantly reduced crying time and salivary cortisol levels.¹⁹³

Conclusions for anesthesia and analgesia for neonatal circumcision:

1. **A DPNB with a ring block, using proper technique, is the most effective technique to provide anesthesia during a neonatal circumcision (Level 1–2, Grade A).**
2. **Topical local anesthetics alone are inferior to nerve and ring blocks and require an adequate time interval for efficacy; they can be used as an adjunct to penile blocks (Level 1–2, Grade A).**
3. **Oral sucrose, non-nutritive sucking, music, and other environmental interventions should only be used as an adjunct to these methods (Level 1–3, Grade A).**

Complications of circumcision

Neonatal circumcision is a safe surgical procedure that is generally well-tolerated. Circumcision complications can



Fig 9. Megameatus intact prepuce hypospadias variant.

vary depending on the surgeon experience, technique used, parental expectation of post-circumcision appearance, timing of circumcision, patient anatomic factors, gestational age, and the accuracy and degree of reporting. Proper pre-operative assessment and examination recognizing possible complicating factors (webbing, ventral skin deficiency, suprapubic fat pad) and adequate postoperative instructions can prevent the commonest complications.

Neonatal circumcisions are performed in the community by a variety of practitioners and the complication rate is higher than that for procedures carried out in the hospital.¹⁹⁴ In addition, it is well-documented that physician ability to self-rate success of a procedure is questionable.¹⁹⁵ When combined with the fact that some of the complications are delayed (e.g., meatal stenosis), there is a high likelihood that the actual complication rate for neonatal circumcisions may be underestimated and constitutes an immense burden to a system where the primary procedure is not covered under universal healthcare.

According to Pieretti et al, almost 5% of pediatric cases performed at a tertiary institution in the U.S. over a five-year span, with an estimated cost of \$685 608, were related to complications of newborn circumcision.¹⁹⁶ The most frequent complications were redundant foreskin and meatal stenosis. Kokorowski et al queried the Pediatric Health Information System database from 28 freestanding pediatric hospitals in the U.S. between 2004 and 2009 and identified those undergoing revision circumcision, non-newborn primary circumcision, and lysis of penile adhesions following neonatal circumcision.¹⁹⁷ The study found a 119% increase in the ratio of revision circumcisions to total male ambulatory procedures during this time period. The median cost for revision circumcision was \$1554 and a total estimated cost of \$6,884,631 for the 28 hospitals over six years.

Overall complication rates

A recent systematic review on complications of neonatal and infant circumcisions noted a wide 0–16% (median 2%) range of adverse events in 16 prospective studies.¹⁹⁸ The same review also found that circumcisions performed in older children were associated with more complications (median 6%) when compared to those carried out in neonates and infants. A systematic review on safety and efficacy of non-therapeutic MC in 5228 men (15–49 years) showed a 4.8% incidence of complications.¹⁹⁹ The most common complication was postoperative infection (1.5%), followed by bleeding (1.3%). Complication rates in the three HIV trials conducted in Africa ranged from 1.7–8%.⁵³⁻⁵⁵

In 1999, the AAP Task Force on Circumcision reported a complication rate of 0.2–0.6%. In contrast, the Canadian Pediatric Society has published complication rates as high as 2% for neonatal circumcision.²⁰⁰ Early complications, such as bleeding and infection, occurred in 0.2% of 136 086 male infants according to a large review.²⁰¹ This study only included complications entered on inpatient records and does not provide an estimate of long-term complications. In another U.S. study of 130 475 newborns conducted in Washington, 0.18% had a bleeding complication after circumcision.²⁰² In a tradeoff analysis, the authors calculated that a complication could be expected in one of every 476 circumcisions, that six UTIs could be prevented for every complication, and about two complications would be expected for every case of penile cancer prevented.²⁰³ A review of complication rates following pediatric circumcision in England between 1997 and 2003 showed that 1.2% of boys experienced a short-term complication and another 0.5% returned to the operating room for a revision within six months.³

Post-circumcision complications can be divided into early and late complications.²⁰⁴⁻²¹¹ Early complications include bleeding, infection, glans necrosis and amputation, delayed/early slippage of circumcision devices, and very rarely death. Late complications include inadequate skin removal, cosmetic issues, inclusion cysts, adhesions and skin bridges, suture sinus tracts, ventral curvature, secondary buried penis and phimosis, urethrocutaneous fistulae, and meatal stenosis.

Factors predicting complication rate

Timing of surgery can be predictive of complications, as bleeding-related complications are higher in older infants.²¹² Penile adhesions and secondary buried penis is more likely in infants with a higher weight for length percentile.²¹³ The results of varying techniques may also be a possible factor affecting complication rates.^{209,214} Results from a small RCT, which compared two surgical techniques (sleeve vs. Plastibell®) in older children, showed that late complications occurred in 12% of the cases that used the sleeve technique

vs. 5% with the Plastibell device.²¹⁴ Current ongoing operator experience was shown to be an important factor in lower bleeding complication rate compared to patient-related variables and long-term operator experience.²¹⁵

Penile adhesions, secondary phimosis, and concealed penis

These relatively common complications of circumcision are the primary reasons for reoperation in the late postoperative period following a circumcision. They are more likely in those with an increased weight for length percentile, a large suprapubic fat pad with abnormal dartos attachments to the skin, or in those with pre-existing peno-scrotal webbing or ventral penile skin deficiency. Topical steroids and occasional dilatation of the prepucial ring and retraction can lead to resolution.^{216, 217} In children with a secondary concealed penis but no phimosis, observation may be an option, as the cosmetic appearance tends to improve with age and surgery should be delayed until the child is at least three years of age.²¹⁸ Adhesions of the mucosal collar to the glans are a common complication and can be prevented by gentle retraction and use of barrier ointments in the early postoperative period.²¹⁹

Meatal stenosis

Meatal stenosis is a common, under-reported complication of circumcision, usually requiring a ventral meatotomy for correction. It is postulated that chronic irritation of the meatus, ammoniacal meatitis, and/or division of the frenular artery after circumcision may be predisposing factors. Meatal stenosis can lead to an upwardly deflected urinary stream, dysuria, urgency, or difficulty with voiding and a flat uroflow curve. The incidence of asymptomatic meatal stenosis (<5 Fr meatal caliber) can be as high as 20%, although its clinical significance is debatable.²⁰⁸ In a review of 1009 circumcised boys who were examined over the age of three years, Van Howe found an overall 2.8% incidence of symptomatic meatal stenosis post-neonatal circumcision.²⁰⁷ Nearly all underwent a meatotomy, but the exact number was unavailable. Studies with long-term followup of circumcision report this complication fairly commonly, with the incidence varying from 2.8–11%.

Recommendations:

- 1. Complication rates post-neonatal circumcision are usually low (around 2%), but given the variability in quoted complication rates and risk of delayed complications not treated by the original physician performing the neonatal circumcision, it is likely that the overall complication rate is slightly higher (Level 2–4).**
- 2. Operator experience and training, recognition of contraindications to circumcision, technique used,**

age, and patient-related variables can impact results and proper reporting and auditing of results is recommended (Level 4, Grade D).

Cost analyses of neonatal circumcision

The potential impact of routine neonatal circumcision needs to be studied from an economic standpoint, as currently, parent-requested circumcision is not covered under most provincial health plans in Canada. The impact on health service use and costs, including direct and indirect procedure costs (costs of managing complications and circumcision revision, litigation costs, and training costs), have to be weighed against indirect cost savings that may be potentially accrued over time, and balanced against the costs of implementing other preventive strategies.

Sansom et al performed a cost effectiveness analysis of newborn circumcision on reducing a U.S. male's lifetime risk of HIV by applying the results of the African trials.²²⁰ The number of circumcisions needed to prevent one HIV infection was 298 for all males, ranging from 65 for black males to 1231 for white males. Newborn circumcision did not generate cost savings for the white males in the U.S., but was a cost saving intervention for all males, African-Americans, and Hispanics. Schoen et al used a third-party U.S. payer database to calculate the cost of newborn circumcision in relation to its health benefits.²²¹ Using a high medically indicated need for post-neonatal circumcision and offsetting costs related to UTI, HIV, balanoposthitis, and penile cancer, the total lifetime net cost of a neonatal circumcision was \$17. The majority of the cost offset (50%) was by the cost of requiring a post-neonatal period circumcision and not the actual health benefits of a circumcision. In a contradictory study by Van Howe, a cost utility analysis did not show circumcision to be cost-effective on sensitivity analysis and on Monte Carlo simulation.²²² A recent study analyzed the impact of a reducing circumcision rate in the U.S. on cost implications related to STIs and UTI.²²³ Reducing circumcision rates from 79% to 10% would increase lifetime direct medical costs by \$313 per foregone circumcision procedure in males.

In a resource-rich setting like Canada, with a relatively lower rate of neonatal circumcisions and an increasing HIV infection risk in MSM, an alternative strategy for MSM males would be a potential strategy. Anderson et al created a mathematical transmission model to conduct an economic analysis and considered four strategies: circumcision of all MSM at 18 years of age, all MSM 35–44 years, all insertive MSM >18 years, and all MSM >18 years.²²⁴ The model predicted a modest 3–5% decrease in HIV infections per year after 25 years of applying these strategies. In the insertive MSM group, 118 circumcisions would need to be performed to prevent one HIV infection. In countries with a high HIV

infection secondary to heterosexual transmission and low circumcision rates, the cost effectiveness of circumcision is more evident.^{225,226} Even in this setting, McAllister et al calculated the NNT to prevent one HIV infection for circumcision as 80 and modeling analysis showed that supplying free condoms was 95 times more cost effective than MC.²²⁷

Training implications

Neonatal circumcision is performed by family physicians, obstetricians, pediatricians, and urologists and this leads to non-standard training methods, varying experience during training and assessment of competency, and varying surgical volumes post-training. Several simulated training methods have been described using penile models to allow standardized teaching using the Mogen and Gomco clamps.^{228,229} In a survey of obstetric-gynecology residents, 63% planned to perform neonatal circumcisions, but only 44% received formal training to do so.²³⁰ When presented with contraindications to circumcision, like hypospadias, buried penis, and micropenis, the average rate of correctly identified contraindications was a dismal 42%. Evidence from the Ugandan trial data showed that approximately 100 procedures are needed to gain competence in sleeve circumcision technique.²³¹ The rate of moderate or severe adverse events showed a statistically significant trend decreasing from 8.8% for the first 20 procedures to around 2% beyond 100. Demaria et al stressed the importance of proper training and followup and the lack thereof of in the current Canadian healthcare system, with unstructured training and inability to deal with contraindications and complications of neonatal circumcision.²³²

Summary of results and recommendations

The effect of MC has to be analyzed at the individual and societal level. For the individual Canadian neonate, there are definite advantages of a circumcision, but the exact estimates of the effect are unknown, the protection provided is not comprehensive, accrue over a lifetime, and can be achieved by other preventive health measures (Table 2). Evidence, therefore, must be analyzed based on its quality and applicability and the GRADE system is an appropriate method to employ when we summarize our results.²³³ There are also clear risks associated with this surgical procedure and parents will continue to have to weigh the potential benefits and risks of neonatal circumcision. In an overall societal perspective, given our healthcare system and the socioeconomic and educational status of our population, universal neonatal circumcision is not justified based on the evidence available.

Competing interests: The authors report no competing personal or financial interests.

Table 2. Benefits of circumcision classified by GRADE recommendations

Clinical benefit	Direction of evidence	Amount of effect	Level of evidence	GRADE quality of evidence	GRADE strength of recommendation
Decreased risk of UTI	Positive	0.07–0.23	Level 2	Low-quality	Weak
Decreased risk of HIV	Positive	0.34–0.62	Level 1	High-quality	Strong*
Decreased risk of HPV prevalence	Positive	0.57–0.77	Level 1	Moderate-quality	Weak
Decreased risk of HPV incidence	Unclear	NS	Level 2	Low-quality	Weak
Decreased risk of HSV	Positive	0.36–0.91	Level 2	Moderate-quality	Weak
Decreased risk of penile cancer	Positive	0.13–0.83	Level 2	Low-quality	Weak

*Concerns related to external validity of data for Canadian population. HPV: human papilloma virus; HSV: herpes simplex virus; UTI: urinary tract infection.

This paper has been peer-reviewed.

References

- Male circumcision. Task Force on Circumcision. *Pediatrics* 2012;130:e756-85. <https://doi.org/10.1542/peds.2012-1990>
- Trends in in-hospital newborn male circumcision-United States. Available at <https://www.cdc.gov/mmwr/preview/mmwrhtml/mm6034a4.htm>. Accessed Feb. 13, 2018.
- Cathcart P, Nuttall M, van der Meulen J, et al. Trends in pediatric circumcision and its complications in England between 1997 and 2003. *Br J Surg* 2006;93:885-90. <https://doi.org/10.1002/bjs.5369>
- Nelson C, Dunn R, Wan J, et al. The increasing incidence of newborn circumcision: Data from the nationwide inpatient sample. *J Urol* 2005;173:978-81. <https://doi.org/10.1097/01.ju.0000145758.80937.7d>
- Wang M, Macklin E, Tracy E, et al. Updated parental viewpoints on male neonatal circumcision in the United States. *Clin Pediatrics* 2010;49:131-6. <https://doi.org/10.1177/0009922809346569>
- Rediger C, Muller AJ. Parent's rationale for male circumcision. *Can Fam Physician* 2013;59:e110-5.
- Centre for Evidence-Based Medicine. Oxford levels of evidence. Available at www.cebm.net. Accessed Jan. 23, 2018.
- Gairdner D. The fate of the foreskin. A study of circumcision. *Br Med J* 1949;2:1433-6. <https://doi.org/10.1136/bmj.2.4642.1433>
- Chang CH, Chang SS. Foreskin development before adolescence in 2149 school-boys. *Int J Urol* 2006;13:968-9. <https://doi.org/10.1111/j.1442-2042.2006.01449.x>
- Babu R, Harrison SK, Hutton KA. Ballooning of the foreskin and physiological phimosis: Is there any evidence of obstructed voiding? *BJU Int* 2004;94:384-7. <https://doi.org/10.1111/j.1464-410X.2004.04935.x>
- McGregor TB, Pike JG, Leonard MP. Pathologic and physiologic phimosis: Approach to the phimotic foreskin. *Can Fam Physician* 2007;53:445-8.
- Letendre J, Barrieras D, Franc-Guimond J, et al. Topical triamcinolone for persistent phimosis. *J Urol* 2009;182:1759-64. <https://doi.org/10.1016/j.juro.2009.03.016>
- de Oliveira Pileggi F, Vincente Y. Phimotic ring topical corticoid cream (0.1% mometasone furoate) treatment in children. *J Pediatr Surg* 2007;42:1749-52. <https://doi.org/10.1016/j.jpedsurg.2007.05.035>
- Lund L, Wai KH, Mui LM, et al. An 18-month followup study after randomized treatment of phimosis in boys with topical steroid vs. placebo. *Scand J Urol Nephrol* 2005;39:78-81. <https://doi.org/10.1080/00365590410002519>
- Nobre YD, Freitas RG, Felizardo MJ, et al. To circ or not to circ: Clinical and pharmaco-economic outcomes of a prospective trial of topical steroid vs. primary circumcision. *Int. Braz J Urol* 2010;36:75-85. <https://doi.org/10.1590/S1677-55382010000100012>
- Yang SS, Tsai YC, Wu CC, et al. Highly potent and moderately potent topical steroids are effective in treating phimosis: A prospective randomized study. *J Urol* 2005;173:1361-3. <https://doi.org/10.1097/01.ju.0000156556.11235.3f>
- Golubovic Z, Milanovic D, Vukadinovic V, et al. The conservative treatment of phimosis in boys. *Br J Urol* 1996;78:786-8. <https://doi.org/10.1046/j.1464-410X.1996.21724.x>
- Zavras N, Christianakis E, Mpourikas D, et al. Conservative treatment of phimosis with fluticasone propionate 0.05%: a clinical study in 1185 boys. *J of Pediatr Urol* 2009;5:181-5. <https://doi.org/10.1016/j.jpuro.2008.11.006>
- Reddy S, Jain V, Dubey M, et al. Local steroid therapy as the first-line treatment for boys with symptomatic phimosis — a long-term prospective study. *Acta Paediatr* 2012;101:130-3. <https://doi.org/10.1111/j.1651-2227.2011.02534.x>
- Ghysel C, Vander Eeck K, Bogaert G. Long-term efficiency of skin stretching and a topical corticoid cream application for unretractable foreskin and phimosis in prepubertal boys. *Urol Int* 2009;82:81-8. <https://doi.org/10.1159/000176031>
- Ku WK, Chiu B, Huen KF. Outcome and recurrence in treatment of phimosis using topical betamethasone in children in Hong Kong. *J Paediatr Child Health* 2007;43:74-9. <https://doi.org/10.1111/j.1440-1754.2007.01006.x>
- Ashfield J, Nickel K, Siemens D, et al. Treatment of phimosis with topical steroids in 194 children. *J Urol* 2003;169:1106-8. <https://doi.org/10.1097/01.ju.0000048973.26072.eb>
- Elmore J, Baker L, Snodgrass W. Tropical steroid therapy as an alternative to circumcision for phimosis in boys younger than 3 years. *J Urol* 2002;168:1746-7. [https://doi.org/10.1016/S0022-5347\(05\)64404-7](https://doi.org/10.1016/S0022-5347(05)64404-7)
- Winberg J, Anderson HJ, Bregstrom T, et al. Epidemiology of symptomatic urinary tract infection in childhood. *Acta Paediatr Scand* 1974;252(suppl):1-20. <https://doi.org/10.1111/j.1651-2227.1974.tb05718.x>
- Panaretto KS, Craig JC, Knight JF, et al. Risk factors for recurrent urinary tract infection in preschool children. *J Paediatr Child Health* 1999;35:454-9. <https://doi.org/10.1046/j.1440-1754.1999.355417.x>
- Shaikh N, Morone N, Bost J et al. Prevalence of urinary tract infection in childhood. *Pediatr Infect Dis J* 2008;27:302-8. <https://doi.org/10.1097/INF.0b013e31815e4122>
- Liu C, Hungate BA, Tobian AR, et al. Male circumcision significantly reduces prevalence and load of genital anaerobic bacteria. *mBio* 2013;4:1-9. <https://doi.org/10.1128/mBio.00076-13>
- Wiswell TE, Miller GM, Gelston HM Jr, et al. Effect of circumcision status on periurethral bacterial flora during the first year of life. *J Pediatr* 1988;113:442-6. [https://doi.org/10.1016/S0022-3476\(88\)80625-5](https://doi.org/10.1016/S0022-3476(88)80625-5)
- Fussell EN, Kaack MB, Cherry R, et al. Adherence of bacteria to human foreskins. *J Urol* 1988;140:997-1001. [https://doi.org/10.1016/S0022-5347\(17\)41909-4](https://doi.org/10.1016/S0022-5347(17)41909-4)
- Balat A, Karakok M, Guler E, et al. Local defense systems in the prepuce. *Scand J Urol Nephrol* 2008;42:63-5. <https://doi.org/10.1080/00365590701472663>
- Ladenhauf HN, Ardelean MA, Schimke C, et al. Reduced bacterial colonization of glans penis after male circumcision in children — a prospective study. *J Pediatr Urol* 2013;9:1137-44. <https://doi.org/10.1016/j.jpuro.2013.04.011>
- Bonacorsi S, Lefevre S, Clermont O, et al. *Escherichia coli* strains causing urinary tract infection in uncircumcised infants resemble urosepsis-like adult strains. *J Urol* 2005;173:195-7. <https://doi.org/10.1097/01.ju.0000148606.07267.7d>
- Shaikh N, Morone N, Lopez J, et al. Does this child have a urinary tract infection? *JAMA* 2007; 298:2895-904. <https://doi.org/10.1001/jama.298.24.2895>
- Singh-Grewal D, Macdessi J, Craig J. Circumcision for the prevention of urinary tract infection in boys: A systematic review of randomized trials and observational studies. *Arch Dis Child* 2005; 90:853-8. <https://doi.org/10.1136/adc.2004.049353>
- Nayir A. Circumcision for the prevention of significant bacteria in boys. *Pediatr Nephrol* 2001;16:1129-34. <https://doi.org/10.1007/s004670100044>
- Morris B, Wiswell T. Circumcision and lifetime risk of urinary tract infection: A systematic review and meta-analysis. *J Urol* 2013;189:2118-24. <https://doi.org/10.1016/j.juro.2012.11.114>
- Jagannath VA, Fedorowicz Z, Sud V, et al. Routine neonatal circumcision for the prevention of urinary tract infections in infancy. *The Cochrane Collaboration* 2012;1-12.
- Zorc J, Levine A, Platt S, et al. Clinical and demographic factors associated with urinary tract infections in young febrile infants. *Pediatrics* 2005;116:644-8. <https://doi.org/10.1542/peds.2004-1825>
- To T, Agha M, Dick P, et al. Cohort study on circumcision of newborn boys and subsequent risk of urinary tract infection. *Lancet* 1998;352:1813-6. [https://doi.org/10.1016/S0140-6736\(98\)02392-7](https://doi.org/10.1016/S0140-6736(98)02392-7)
- Craig J, Knight J, Sureshkumar P, et al. Effect of circumcision on incidence of urinary tract infection in preschool boys. *J Pediatr* 1996;128: 23-7. [https://doi.org/10.1016/S0022-3476\(96\)70423-7](https://doi.org/10.1016/S0022-3476(96)70423-7)
- Simforosh N, Tabibi A, Khalili, S, et al. Neonatal circumcision reduces the incidence of asymptomatic urinary tract infection: A large prospective study with long-term followup using Plastibell. *J Pediatr Urol* 2012;8:320-3. <https://doi.org/10.1016/j.jpuro.2010.10.008>
- Toker O, Schwartz S, Segal G, et al. A costly covenant: Ritual circumcision and urinary tract infection. *IMAJ* 2010;12:262-5.
- Prais D, Shoo-Furman R, Amir J. Is ritual circumcision a risk factor for neonatal urinary tract infections? *Arch Dis Child* 2009;94:191-4. <https://doi.org/10.1136/adc.2008.144063>

44. Alsaywid B, Saleh H, Deshpande A, et al. High-grade primary vesicoureteral reflux in boys: Long-term results of a prospective cohort study. *J Urol* 2010;184:1598-1603. <https://doi.org/10.1016/j.juro.2010.04.021>
45. Kwak C, Seung-June OH, Lee A, et al. Effect of circumcision on urinary tract infection after successful antireflux surgery. *BJU Int* 2004;94:627-9. <https://doi.org/10.1111/j.1464-410X.2004.05014.x>
46. Gucuk A, Burgu B, Gokce L, et al. Do antibiotic prophylaxis and/or circumcision change periurethral uropathogen colonization and urinary tract infection rates in boys with VUR? *J Pediatr* 2013;1-6.
47. Mukherjee S, Joshi A, Carroll D, et al. What is the effect of circumcision on risk of urinary tract infection in boys with posterior urethral valves? *J Pediatr* 2009 44:417-21.
48. Zareba P, Lorenzo AJ, Braga LH. Risk factors for febrile urinary tract infection in infants with prenatal hydronephrosis: Comprehensive, single-centre analysis. *J Urol* 2014;191:1614-8. <https://doi.org/10.1016/j.juro.2013.10.035>
49. Roth C, Hubanks M, Bright B, et al. Occurrence of urinary tract infection in children with significant upper urinary tract obstruction. *J Urol* 2009;73:74-8. <https://doi.org/10.1016/j.urol.2008.05.021>
50. Song SH, Lee SB, Park YS, et al. Is antibiotic prophylaxis necessary in infants with obstructive hydronephrosis? *J Urol* 2007;177:1098-101. <https://doi.org/10.1016/j.juro.2006.11.002>
51. Wingerter S, Bachur R. Risk factors for contamination of catheterized urine specimens in febrile children. *Pediatr Emerg Care* 2011;27:1-4. <https://doi.org/10.1097/PEC.0b013e3182037c20>
52. Infectious Disease Prevention and Control: Estimates of HIV prevalence and incidence in Canada, 2011. Available at https://www.canada.ca/content/dam/phac-aspc/migration/phac-aspc/aids-sida/publication/epi/2010/pdf/EN_Chapter1_Web.pdf. Accessed December 19, 2017.
53. Bailey R, Moses S, Parker C, et al. Male circumcision for HIV prevention in young men in Kisumu, Kenya: A randomized controlled trial. *Lancet* 2007;369:643-56. [https://doi.org/10.1016/S0140-6736\(07\)60312-2](https://doi.org/10.1016/S0140-6736(07)60312-2)
54. Gray R, Kigozi G, Serwadda D, et al. Male circumcision for HIV prevention in med in Rakai, Uganda: A randomized trial. *Lancet* 2007;369:657-66. [https://doi.org/10.1016/S0140-6736\(07\)60313-4](https://doi.org/10.1016/S0140-6736(07)60313-4)
55. Auvert B, Taljaard D, Lagarde E, et al. Randomized, controlled intervention trial of male circumcision for reduction of HIV infection risk: The ANRS 1265 Trial. *PLoS Medicine* 2005;2:1112-22. <https://doi.org/10.1371/journal.pmed.0020298>
56. WHO/UNAIDS. New data on male circumcision and HIV prevention: policy and programme implications. Geneva: WHO; 2007.
57. Smith D, Taylor A, Kilmarx P, et al. Male circumcision in the United States for the prevention of HIV infection and other adverse health outcomes: Report from a CDC consultation. *Public Health Records* 2010;125:72-82. <https://doi.org/10.1177/00335491012505110>
58. Mehta SD, Krieger JN, Agot K, et al. Circumcision and reduced risk of self-reported penile coital injuries: Results from a randomized controlled trial in Kisumu, Kenya. *J Urol* 2010;184:203-7. <https://doi.org/10.1016/j.juro.2010.03.015>
59. Soto-Ramirez LE, Renjifo B, McLane MF, et al. HIV-1 Langerhans' cell tropism associated with heterosexual transmission of HIV. *Science* 1996;271:1291-3. <https://doi.org/10.1126/science.271.5253.1291>
60. Price LB, Liu CM, Johnson KE, et al. The effects of circumcision on the penis microbiome. *PLoS One* 2010;5:e8422. <https://doi.org/10.1371/journal.pone.0008422>
61. McCoombe SG, Short RV. Potential HIV-1 target cells in the human penis. *AIDS* 2006;20:1491-5. <https://doi.org/10.1097/01.aids.0000237364.11123.98>
62. Kigozi G, Wawer M, Ssetuba A, et al. Foreskin surface area and HIV acquisition in Rakai, Uganda (size matters). *AIDS* 2009;23:2209-13. <https://doi.org/10.1097/QAD.0b013e3283303eda8>
63. Sobngwi-Tambekou J, Taljaard D, Lissouba P, et al. Effect of HSV-2 Serostatus on acquisition of HIV by young men: Results of a longitudinal study in Orange Farm, South Africa. *J Infect Dis* 2009;199:958-63. <https://doi.org/10.1086/597208>
64. Gray R, Serwadda D, Tobian A, et al. Effects of genital ulcer disease and herpes simplex virus type 2 on the efficacy of male circumcision for HIV prevention: Analyses from the Rakai trials. *PLoS* 2009;6:1-8. <https://doi.org/10.1371/journal.pmed.1000187>
65. Mehta S, Moses S, Parker C, et al. Circumcision status and incident herpes simplex virus type 2 infection, genital ulcer disease, and HIV infection. *AIDS* 2012;26:1141-9. <https://doi.org/10.1097/QAD.0b013e328352d116>
66. Sahasrabudhe V, Vermund S. The future of HIV prevention: Control of sexually transmitted infections and circumcision interventions. *Infect Dis Clin Am* 2007;21:241-57. <https://doi.org/10.1016/j.idc.2007.03.005>
67. Fink AJ. A possible explanation for heterosexual male infection with AIDS. *N Engl J Med* 1986;315:1167. <https://doi.org/10.1056/NEJM198610303151818>
68. Weiss HA, Quigley MA, Hayes RJ. Male circumcision and risk of HIV infection in sub-Saharan Africa: A systematic review and meta-analysis. *AIDS* 2000;14: 2361-70. <https://doi.org/10.1097/00002030-200010200-00018>
69. Warner L, Ghanem K, Newman D, et al. Male circumcision and risk of HIV infection among heterosexual African American men attending Baltimore sexually transmitted disease clinics. *J Infect Dis* 2009;199:65. <https://doi.org/10.1086/595569>
70. Drain PK, Halperin DT, Hughes JP, et al. Male circumcision, religion, and infectious diseases: An ecologic analysis of 118 developing countries. *BMC Infect Dis* 2006;6:172. <https://doi.org/10.1186/1471-2334-6-172>
71. Siegfried N, Muller M, Deeks JJ, et al. Male circumcision for prevention of heterosexual acquisition of HIV in men (review). *The Cochrane Library* 2009;4:1-40. <https://doi.org/10.1002/14651858.CD003362.pub2>
72. Eaton L, Cain D, Agrawal A, et al. The influence of male circumcision for HIV prevention on sexual behaviour among traditionally circumcised men in Cape Town, South Africa. *Int J STD AIDS* 2011;22:6674-9.
73. Gray R, Kigozi G, Kong X, et al. The effectiveness of male circumcision for HIV prevention and effects on risk behaviours in a post-trial followup study. *AIDS* 2012;5:609-15. <https://doi.org/10.1097/QAD.0b013e3283504a3f>
74. Mattson C, Campbell R, Bailey R, et al. Risk compensation is not associated with male circumcision in Kisumu, Kenya: A multi-faceted assessment of men enrolled in a randomized controlled trial. *PLoS* 2008;3:1-9. <https://doi.org/10.1371/journal.pone.0002443>
75. Hall HI, Song R, Kaplan EH, et al. Estimation of HIV incidence in the United States. *JAMA* 2008;300:520-9. <https://doi.org/10.1001/jama.300.5.520>
76. Wiysonge CS, Kongnyuy EJ, Shey M, et al. Male circumcision for prevention of homosexual acquisition of HIV in men. *Cochrane Database Syst Rev* 2011;(6):CD007496. <https://doi.org/10.1002/14651858.CD007496.pub2>
77. Millett GA, Flores SA, Marks G, et al. Circumcision status and risk of HIV and sexually transmitted infections among men who have sex with men: A meta-analysis. *JAMA* 2008; 300:1674-84. <https://doi.org/10.1001/jama.300.14.1674>
78. Londish GJ, Templeton DJ, Regan DJ, et al. Minimal impact of circumcision on HIV acquisition in men who have sex with men. *Sex Health* 2010;7:463-70. <https://doi.org/10.1071/SH09080>
79. Wawer M, Makumbi F, Kigozi G, et al. Circumcision in HIV-infected men and its effect on HIV transmission to female partners in Rakai, Uganda: A randomized controlled trial. *Lancet* 2009;374:229-37. [https://doi.org/10.1016/S0140-6736\(09\)60998-3](https://doi.org/10.1016/S0140-6736(09)60998-3)
80. Weiss H, Hankins C, Dickson K. Male circumcision and risk of HIV infection in women: A systematic review and meta-analysis. *Lancet* 2009;9:669-77. [https://doi.org/10.1016/S1473-3099\(09\)70235-X](https://doi.org/10.1016/S1473-3099(09)70235-X)
81. Baeten J, Donnell D, Kapinga S, et al. Male circumcision and risk of male-to-female HIV-1 transmission: A multinational prospective study in African HIV-1 serodiscordant couples. *AIDS* 2010;13:1-13. <https://doi.org/10.1097/QAD.0b013e32833616e0>
82. Green L, Travis J, McAllister R, et al. Male circumcision and HIV prevention insufficient evidence and neglected external validity. *Am J Prev Med* 2010;39:479-82. <https://doi.org/10.1016/j.amepre.2010.07.010>
83. Newell M-L, Barnighausen T. Male circumcision to cut HIV risk in the general population. *Lancet* 2007;369:617-9. [https://doi.org/10.1016/S0140-6736\(07\)60288-8](https://doi.org/10.1016/S0140-6736(07)60288-8)
84. Lie R, Miller F. What counts as reliable evidence for public health policy: The case of circumcision for preventing HIV infection. *BMC Med Res Methodol* 2011;11:1-7. <https://doi.org/10.1186/1471-2288-11-34>
85. Weiss H, Dickson K, Agot K, et al. Male circumcision for HIV prevention: Current research and programmatic issues. *AIDS* 2010;24:S61-8. <https://doi.org/10.1097/01.aids.0000390708.66136.f4>
86. Mor Z, Kent C, Kohn R, et al. Declining rates in male circumcision amidst increasing evidence of its public health benefit. *PLoS* 2007;9:1-4. <https://doi.org/10.1371/journal.pone.0000861>
87. Xu X, Patel D, Dalton V, et al. Can routine neonatal circumcision help prevent human immunodeficiency virus transmission in the United States? *Am J Mens Health* 2009;3:79-84. <https://doi.org/10.1177/1557988308323616>
88. Auvert B, Sobngwi-Tambekou J, Cutler E, et al. Effect of male circumcision on the prevalence of high-risk human papillomavirus in young men: Results of a randomized controlled trial conducted in Orange Farm, South Africa. *J Infect Dis* 2009;199:14-19. <https://doi.org/10.1086/595566>
89. Tobian A, Serwadda D, Quinn T, et al. Male circumcision for the prevention of HSV-2 and HPV infections and syphilis. *N Engl J Med* 2009;13:1298-1309. <https://doi.org/10.1056/NEJMoa0802556>
90. Tobian A, Kong X, Gravitt P, et al. Male circumcision and anatomic sites of penile high-risk human papillomavirus in Rakai, Uganda. *Int J Cancer* 2011;12:1-11. <https://doi.org/10.1002/ijc.25957>
91. Gray R, Serwadda D, Kong X. Male circumcision decreases acquisition and increases clearance of high-risk human papillomavirus in HIV-negative men: A randomized trial in Rakai, Uganda. *J Infect Dis* 2010;15:1-14. <https://doi.org/10.1086/652184>
92. Serwadda D, Wawer M, Makumbi F, et al. Circumcision of HIV-infected men: Effects on high-risk human papillomavirus infections in a randomized trial in Rakai, Uganda. *J Infect Dis* 2010;201:1-14. <https://doi.org/10.1086/652185>
93. Tobian AA, Kigozi G, Gravitt PE, et al. Human papillomavirus incidence and clearance among HIV-positive and HIV-negative men in sub-Saharan Africa. *AIDS* 2012;26:1555-65. <https://doi.org/10.1097/QAD.0b013e328353b83c>
94. Wilson L, Gravitt P, Tobian A et al. Male circumcision reduces penile high-risk human papillomavirus viral load in a randomized clinical trial in Rakai, Uganda. *Sex Transm Infect* 2013;89:262-6. <https://doi.org/10.1136/sextrans-2012-050633>
95. Weaver BA, Feng Q, Holmes KK, et al. Evaluation of genital sites and sampling techniques for detection of human papillomavirus DNA in men. *J Infect Dis* 2004;189:677-85. <https://doi.org/10.1086/381395>

96. Larke N, Thomas SL, Dos Santos S, et al. Male circumcision and human papillomavirus infection in men: A systematic review and meta-analysis. *J Infect Dis* 2011;9:1375-90. <https://doi.org/10.1093/infdis/jir523>
97. Albero G, Castellsague X, Giuliano A, et al. Male circumcision and genital human papillomavirus: A systematic review and meta-analysis. *Sex Transm Dis* 2012;39:104-13. <https://doi.org/10.1097/OLQ.0b013e3182387abd>
98. Dickson N, Ryding J, van Roode T, et al. Male circumcision and serologically determined human papillomavirus infection in a birth cohort. *Cancer Epidemiol Biomarkers* 2009;18:177-83. <https://doi.org/10.1158/1055-9965.EPI-08-0353>
99. Albero G, Villa L, Lazzano-Ponce E, et al. Male circumcision and prevalence of genital human papillomavirus infection in men? A multinational study. *BMC Infect Dis* 2013;13:1-10. <https://doi.org/10.1186/1471-2334-13-18>
100. Poynten I, Jin F, Templeton D, et al. Prevalence, incidence, and risk factors for human papillomavirus 16 seropositivity in Australian homosexual men. *Sex Transm Dis* 2012;39: 726-32. <https://doi.org/10.1097/OLQ.0b013e31825d5cb8>
101. Wawer M, Tobian A, Kigozi G, et al. Effect of circumcision of HIV-negative men on transmission of human papillomavirus to HIV-negative women: A randomized trial in Rakai, Uganda. *The Lancet* 2011; 377: 209-18. [https://doi.org/10.1016/S0140-6736\(10\)61967-8](https://doi.org/10.1016/S0140-6736(10)61967-8)
102. Tobian A, Kong X, Wawer M, et al. The effect of circumcision of HIV-infected men on human papillomavirus infection in female partners: Analyses using data from a randomized trial in Rakai, Uganda. *Lancet Infect Dis* 2011;8:604-12. [https://doi.org/10.1016/S1473-3099\(11\)70038-X](https://doi.org/10.1016/S1473-3099(11)70038-X)
103. Winer RL, Hughes JP, Feng Q, et al. Condom use and the risk of genital human papillomavirus infection in young women. *N Engl J Med* 2006;354: 2645-54. <https://doi.org/10.1056/NEJMoa053284>
104. Mehta S, Moses S, Agot K, et al. Adult male circumcision does not reduce the risk of incident *Neisseria gonorrhoeae*, *Chlamydia trachomatis*, or *Trichomonas vaginalis* infection: Results from a randomized, controlled trial in Kenya. *J Infect Dis* 2009;8:370-8. <https://doi.org/10.1086/600074>
105. Sobngwi-Tambekou J, Tijaard D, Nieuwoudt M, et al. Male circumcision and *Neisseria gonorrhoeae*, *Chlamydia trachomatis*, and *Trichomonas vaginalis*: Observations after a randomized controlled trial for HIV prevention. *Sex Transm Infect* 2009;85:116-20. <https://doi.org/10.1136/sti.2008.032334>
106. Weiss HA, Thomas SL, Munabi SK, et al. Male circumcision and risk of syphilis, chancroid, and genital herpes: A systematic review and meta-analysis. *Sex Transm Infect* 2006;82:101-10. <https://doi.org/10.1136/sti.2005.017442>
107. Disaker R, Peterman T, Kamb M, et al. Circumcision and STD in the United States: Cross-sectional and cohort analyses. *Sex Transm Inf* 2000;474-9.
108. Sena AC, Miller WC, Hobbs MM, et al. *Trichomonas vaginalis* infection in male sexual partners: Implications for diagnosis, treatment, and prevention. *Clin Infect Dis* 2007;44:13-22. <https://doi.org/10.1086/511144>
109. Mehta S, Gaydos C, Macean I, et al. The effect of medical male circumcision on urogenital *Mycoplasma genitalium* among men in Kisumu, Kenya. *Sex Transm Dis* 2012;39:276-80. <https://doi.org/10.1097/OLQ.0b013e318240189c>
110. Turner A, Morrison C, Padian N, et al. Male circumcision and women's risk of incident chlamydial, gonococcal, and trichomonal infections. *Sex Transm Dis* 2008;37:689-95. <https://doi.org/10.1097/OLQ.0b013e31816b1fcc>
111. Tobian A, Quinn T. Herpes simplex virus type 2 and syphilis infections with HIV: An evolving synergy in transmission and prevention. *Curr Opin HIVAIDS* 2009;4:294-9. <https://doi.org/10.1097/COH.0b013e3182832c1881>
112. Powers KA, Poole C, Pettifor AE, et al. Re-thinking the heterosexual infectivity of HIV-1: A systematic review and meta-analysis. *Lancet Infect Dis* 2008;8:553-63. [https://doi.org/10.1016/S1473-3099\(08\)70156-7](https://doi.org/10.1016/S1473-3099(08)70156-7)
113. Freeman EE, Weiss HA, Glynn JR, et al. Herpes simplex virus 2 infection increases HIV acquisition in men and women: Systematic review and meta-analysis of longitudinal studies. *AIDS* 2006;20:73-83. <https://doi.org/10.1097/01.aids.0000198081.09337.a7>
114. Barnabas RV, Wasserheit JN, Huang Y, et al. Impact of herpes simplex virus Type-2 on HIV-1 acquisition and progression in an HIV vaccine trial (the Step Study). *J Acquir Immune Defic Syndr* 2011;57:238-44. <https://doi.org/10.1097/QAI.0b013e31821acb5>
115. Watson-Jones D, Weiss HA, Rusizoka M, et al. Effect of herpes simplex suppression on incidence of HIV among women in Tanzania. *N Engl J Med* 2008;358:1560-71. <https://doi.org/10.1056/NEJMoa0800260>
116. Celum C, Wald A, Hughes J, et al. Effect of acyclovir on HIV-1 acquisition in herpes simplex virus 2 seropositive women and men who have sex with men: A randomized, double-blind, placebo-controlled trial. *Lancet* 2008;371:2109-19. [https://doi.org/10.1016/S0140-6736\(08\)60920-4](https://doi.org/10.1016/S0140-6736(08)60920-4)
117. Tobian A, Ssempiija V, Kigozi G, et al. Incident HIV and herpes simplex virus type 2 infection among men in Rakai, Uganda. *AIDS* 2009;23:1589-94. <https://doi.org/10.1097/QAD.0b013e3182832d4042>
118. Tobian A, Charvat B, Ssempiija V, et al. Factors associated with the prevalence and incidence of herpes simplex virus Type-2 infection among men in Rakai, Uganda. *J Infect Dis* 2009;199:945-9. <https://doi.org/10.1086/597074>
119. Xu F, Markowitz L, Sternberg M, et al. Prevalence of circumcision and herpes simplex virus Type-2 infection in men in the United States: The national health and nutrition examination survey (NHANES), 1999–2004. *Sex Transm Dis* 2007;34:479-84. <https://doi.org/10.1097/01.olq.0000253335.41841.04>
120. Jameson D, Celum C, Manhart L, et al. The association between lack of circumcision and HIV, HSV-2 and other sexually transmitted infections among men who have sex with men. *Sex Transm Dis* 2010;37:147-52. <https://doi.org/10.1097/OLQ.0b013e3181bd0ff0>
121. Dickson N, van Roode T, Herbison P, et al. Circumcision and risk of sexually transmitted infections in a birth cohort. *Sex Transm Inf* 2008;383-7.
122. Fergusson D, Boden J, Horwood J. Circumcision status and risk of sexually transmitted infection in young adult males: An analysis of longitudinal birth cohort. *Pediatrics* 2006;118:1971-7. <https://doi.org/10.1542/peds.2006-1175>
123. Gray R, Kigozi G, Serwadda D, et al. The effects of male circumcision on female partners' genital tract symptoms and vaginal infections in a randomized trial in Rakai, Uganda. *Am J Obstet Gynecol* 2009;42:1-42.7.
124. Tobian AA, Kigozi G, Redd AD, et al. Male circumcision and herpes simplex virus type 2 infection in female partners: A randomized trial in Rakai, Uganda. *J Infect Dis* 2012;205:486-90. <https://doi.org/10.1093/infdis/jir767>
125. Jemal A, Siegel R, Ward E, et al. Cancer statistics, 2007. *CA Cancer J Clin* 2007;57:43-66. <https://doi.org/10.3322/canjclin.57.1.43>
126. Pow-Sang MR, Ferreira U, Pow-Sang JM, et al. Epidemiology and natural history of penile cancer. *Urology* 2010;76:S2-6.
127. Bleeker M, Heideman D, Snijders P, et al. Penile cancer: Epidemiology, pathogenesis, and prevention. *World J Urol* 2009;27:141-50. <https://doi.org/10.1007/s00345-008-0302-z>
128. Minhas S, Manseck A, Watya, et al. Penile cancer — prevention and premalignant conditions. *Urology* 2010;76:S24-35. <https://doi.org/10.1016/j.urol.2010.04.007>
129. Shabbir M, Barod R, Hegarty P, et al. Primary prevention and vaccination for penile cancer. *Ther Adv Urol* 2013;5:161-9. <https://doi.org/10.1177/1756287212465456>
130. Miralles-Guri C, Bruni L, Cubilla AL, et al. Human papillomavirus prevalence and type distribution in penile carcinoma. *J Clin Pathol* 2009;62:870-8. <https://doi.org/10.1136/jcp.2008.063149>
131. Dillner J, von Krogh G, Horenblas S, et al. Etiology of squamous cell carcinoma of the penis. *Scand J Urol Nephrol* 2000;205:189-93. <https://doi.org/10.1080/00365590050509913>
132. Parkin DM, Bray F. Chapter 2: The burden of HPV-related cancers. *Vaccine* 2006;24:11-25. <https://doi.org/10.1016/j.vaccine.2006.05.111>
133. Larke N, Thomas SL, dos Santos Silva I, et al. Male circumcision and penile cancer: A systematic review and meta-analysis. *Cancer Causes Control* 2011;22:1097-1110. <https://doi.org/10.1007/s10552-011-9785-9>
134. D'Hauwers KWM, Depuydt CE, Bogers JJ, et al. Human papillomavirus, lichen sclerosus, and penile cancer: A study in Belgium. *Vaccine* 2012;30:6573-7. <https://doi.org/10.1016/j.vaccine.2012.08.034>
135. Tsen HF, Morgenstern H, Mack T, et al. Risk factors for penile cancer: Results of a population-based case-control study in Los Angeles County (United States). *Cancer Causes Control* 2001;12:267-77. <https://doi.org/10.1023/A:1011266405062>
136. Madsen BS, van den Brule AJC, Jensen HL, et al. Risk factors for squamous cell carcinoma of the penis — population-based case-control study in Denmark. *Cancer Epidemiol Biomarkers* 2008;17:2683-91. <https://doi.org/10.1158/1055-9965.EPI-08-0456>
137. Daling J, Madeline M, Johnson L, et al. Penile cancer: Importance of circumcision, human papillomavirus, and smoking in situ and invasive disease. *Int J Cancer* 2005;116:606-16. <https://doi.org/10.1002/ijc.21009>
138. Frisch M, Friis S, Kjaer SK, et al. Falling incidence of penis cancer in an uncircumcised population (Denmark 1943-90). *BMJ* 1995;311:1471. <https://doi.org/10.1136/bmj.311.7018.1471>
139. Baldur-Felskov B, Hannibal CG, Mung C, et al. Increased incidence of penile cancer and high-grade penile intraepithelial neoplasia in Denmark 1978–2008: A nationwide population-based study. *Cancer Causes Control* 2012;23:273-80. <https://doi.org/10.1007/s10552-011-9876-7>
140. Goodman M, Hernandez B, Shvesov Y. Demographic and pathologic differences in the incidence of invasive penile cancer in the United States, 1995–2003. *Cancer Epidemiol Biomarkers* 2007;16:1833-9. <https://doi.org/10.1158/1055-9965.EPI-07-0221>
141. Taylor ML, Mainous AG 3rd, Wells BJ. Prostate cancer and sexually transmitted diseases: A meta-analysis. *Fam Med* 2005;37:506-12.
142. Wright J, Lin D, Stanford J. Circumcision and the risk of prostate cancer. *Cancer* 2011;118:4437-43. <https://doi.org/10.1002/cncr.26653>
143. Ewing P, Bowie C. A case-control study of cancer of the prostate in Somerset and east Devon. *Br J Cancer* 1996;74:661-6. <https://doi.org/10.1038/bjc.1996.418>

144. Tian Y, Liu W, Wang JZ, et al. Effects of circumcision on male sexual functions: A systematic review and meta-analysis. *Asian J Androl* 2013;15:662-6. <https://doi.org/10.1038/aja.2013.47>
145. Kigozi G, Watya S, Polist C, et al. The effect of male circumcision on sexual satisfaction and function, results from a randomized trial of male circumcision for human immunodeficiency virus prevention, Rakai, Uganda. *BJU Int* 2008;101:65-70. <https://doi.org/10.1111/j.1464-410X.2007.07369.x>
146. Krieger J, Mehta S, Bailey R, et al. Adult male circumcision: Effects on sexual function and sexual satisfaction in Kisumu, Kenya. *J Sex Med* 2008;5:2610-22. <https://doi.org/10.1111/j.1743-6109.2008.00979.x>
147. Mehta S, Krieger J, Agot K, et al. Circumcision and reduced risk of self-reported penile coital injuries: Results from a randomized controlled trial in Kisumu, Kenya. *J Urol* 2010;184:203-7. <https://doi.org/10.1016/j.juro.2010.03.015>
148. Frisch M, Lindholm M, Gronbaek M. Male circumcision and sexual function in men and women: A survey-based, cross-sectional study in Denmark. *Int J Epidemiol* 2011;40:1367-81. <https://doi.org/10.1093/ije/dyr104>
149. Bronselaer G, Schober J, Meyer-Bahlburg H, et al. Male circumcision decreases penile sensitivity as measured in a large cohort. *BJU Int* 2013;111:820-7. <https://doi.org/10.1111/j.1464-410X.2012.11761.x>
150. Sorrells M, Snyder J, Reiss M, et al. Fine-touch pressure thresholds in the adult penis. *BJU Int* 2007;99:864-9. <https://doi.org/10.1111/j.1464-410X.2006.06685.x>
151. Kim D, Pang MG. The effect of male circumcision on sexuality. *BJU Int* 2006;99:619-22. <https://doi.org/10.1111/j.1464-410X.2006.06646.x>
152. Collins S, Upshaw J, Rutchik S, et al. Effects of circumcision on male sexual function: Debunking a myth? *J Urol* 2002;167:2111-2. [https://doi.org/10.1016/S0022-5347\(05\)65097-5](https://doi.org/10.1016/S0022-5347(05)65097-5)
153. Peterson A. Elective, adult circumcision does not affect patient perception of sexual health as defined by the Male Sexual Questionnaire (MSHQ). *JMH* 2010;7:368-72.
154. Senkul T, Iseri C, Sen B, et al. Circumcision in adults: Effect on sexual function. *J Urol* 2004;63:155-8. <https://doi.org/10.1016/j.urolgy.2003.08.035>
155. Waldinger M, Quinn P, Dilleen M, et al. A multinational population survey of intravaginal ejaculation latency time. *J Sex Med* 2005;2:492-7. <https://doi.org/10.1111/j.1743-6109.2005.00070.x>
156. Waldinger M, McIntosh J, Schweitzer D. A five-nation survey to assess the distribution of the intravaginal ejaculatory latency time among the general male population. *J Sex Med* 2009;6:2888-95. <https://doi.org/10.1111/j.1743-6109.2009.01392.x>
157. Bleustein C, Fogarty J, Eckholdt H, et al. Effect of neonatal circumcision on penile neurologic sensation. *J Urol* 2005;65:773-7. <https://doi.org/10.1016/j.urolgy.2004.11.007>
158. Kigozi G, Lukabwale I, Kagaayi J, et al. Sexual satisfaction of women partners of circumcised men in randomized trial of male circumcision in Rakai, Uganda. *BJU Int* 2009;104:1698-1701. <https://doi.org/10.1111/j.1464-410X.2009.08683.x>
159. Mao L, Templeton DJ, Crawford J, et al. Does circumcision make a difference to the sexual experience of gay men? Findings from the Health in Men (HiM) cohort. *J Sex Med* 2008;5:2557-61. <https://doi.org/10.1111/j.1743-6109.2008.00845.x>
160. Shankar KR, Rickwood AM. The incidence of phimosis in boys. *BJU Int* 1999;84:101-2. <https://doi.org/10.1046/j.1464-410x.1999.00147.x>
161. McGregor T, Pike J, Leonard M. Pathologic and physiologic phimosis: Approach to the phimotic foreskin. *Can Fam Physician* 2007;53:445-8.
162. Metcalfe P. Foreskin management. *Can Fam Physician* 2010;56:290-5.
163. Kayaba H, Tamura H, Kitajima S, et al. Analysis of shape and retractability of the prepuce in 603 Japanese boys. *J Urol* 1996;156:1813-5. [https://doi.org/10.1016/S0022-5347\(01\)65544-7](https://doi.org/10.1016/S0022-5347(01)65544-7)
164. Malone P, Steinbrecher H. Medical aspects of male circumcision. *BMJ* 2007;335:1206-90. <https://doi.org/10.1136/bmj.39385.382708.AD>
165. Rickwood AMK. Medical indications for circumcision. *BJU Int* 1999;83:45-51. <https://doi.org/10.1046/j.1464-410x.1999.0830s1045.x>
166. Hutcheson J. Male neonatal circumcision: Indications, controversies, and complications. *Urol Clin N Am* 2004;31:461-7. <https://doi.org/10.1016/j.ucl.2004.04.011>
167. Clouston D, Hall A, Lawrenschuk N. Penile lichen sclerosus (balanitis xerotica obliterans). *BJU Int* 2011;108:14-9. <https://doi.org/10.1111/j.1464-410X.2011.10699.x>
168. Gargallo P, Kozakewich H, Bauer S, et al. Balanitis xerotica obliterans in boys. *J Urol* 2005;174:1409-12. <https://doi.org/10.1097/01.ju.0000173126.63094.b3>
169. Becker K. Lichen Schlerosus in boys. *Dtsch Arztebl* 2011;108:53-8.
170. Kiss A, Kiraly L, Kutasy B, et al. High incidence of balanitis xerotica obliterans in boys with phimosis: Prospective 10-year study. *Ped Dermatol* 2005;22:305-8. <https://doi.org/10.1111/j.1525-1470.2005.22404.x>
171. Meuli M, Briner J, Hanimann B, et al. Lichen sclerosus et atrophicus causing phimosis in boys: A prospective study with 5-year followup after complete circumcision. *J Urol* 1994;152:987-9. [https://doi.org/10.1016/S0022-5347\(17\)32638-1](https://doi.org/10.1016/S0022-5347(17)32638-1)
172. Mattioli G, Repetto P, Carlini C, et al. Lichen sclerosus et atrophicus in children with phimosis and hypospadias. *Pediatr Surg Int* 2002;18:273-5. <https://doi.org/10.1007/s003830100699>
173. Yardley IE, Cosgrove C, Lambert AW. Pediatric preputial pathology: Are we circumcising enough? *Ann R Coll Surg Engl* 2007;89:62-5. <https://doi.org/10.1308/003588407X160828>
174. Bochove-Overgaauw DM, Gelders W, de Vylder AMA. Routine biopsies in pediatric circumcision: (Non) sense? *J Pediatr Urol* 2009;5:178-80. <https://doi.org/10.1016/j.jpuro.2008.11.008>
175. Vincent MV, Mackinnon E. The response of clinical balanitis xerotica obliterans to the application of topical steroid-based creams. *J Pediatr Surg* 2005;40:709-12. <https://doi.org/10.1016/j.jpedsurg.2004.12.001>
176. Holbrook C, Tsang T. Management of boys with abnormal appearance of meatus at circumcision for balanitis xerotica obliterans. *Ann R Coll Surg Engl* 2011;93:482-4. <https://doi.org/10.1308/003588411X587145>
177. Snodgrass W, Khavari R. Prior circumcision does not complicate the repair of hypospadias with an intact prepuce. *J Urol* 2006;176:296-8. [https://doi.org/10.1016/S0022-5347\(06\)00564-7](https://doi.org/10.1016/S0022-5347(06)00564-7)
178. Rodriguez V, Titapiwatanakun R, Moir C, et al. To circumcise or not to circumcise? Circumcision in patients with bleeding disorders. *Haemophilia* 2009;16:272-6. <https://doi.org/10.1111/j.1365-2516.2009.02119.x>
179. Kenet G, Chan AKC, Soucie JM, et al. Bleeding disorders in neonates. *Haemophilia* 2010;16:168-75. <https://doi.org/10.1111/j.1365-2516.2010.02316.x>
180. Brady-Fryer B, Wiebe N, Lander J. Pain relief for neonatal circumcision. *Intervention Review* 2009;1-113.
181. Lehr V, Taddio A. Topical anesthesia in neonates: Clinical practices and practical considerations. *Semin Perinatol* 2007;31:323-9. <https://doi.org/10.1053/j.semper.2007.07.008>
182. Taddio A, Goldbach M, Ipp M, et al. Effect of neonatal circumcision on pain responses during vaccination in boys. *Lancet* 1995;345:291-2. [https://doi.org/10.1016/S0140-6736\(95\)90278-3](https://doi.org/10.1016/S0140-6736(95)90278-3)
183. Banihghbal B. Optimal time for neonatal circumcision: An observation-based study. *J Pediatr Urol* 2009;5:359-62. <https://doi.org/10.1016/j.jpuro.2009.01.002>
184. Taddio A, Stevens B, Craig K, et al. Efficacy and safety of lidocaine-prilocaine cream for pain during circumcision. *N Engl J Med* 1997;336:1197-201. <https://doi.org/10.1056/NEJM199704243361701>
185. Wahlgren CF, Quiding H. Depth of cutaneous analgesia after application of a eutectic mixture of the local anesthetics lidocaine and prilocaine (EMLA cream). *J Am Acad Dermatol* 2000;42:584-8. <https://doi.org/10.1067/mjd.2000.104303>
186. Lander J, Brady-Fryer B, Metcalfe JB, et al. Comparison of ring block, dorsal penile nerve block, and topical anesthesia for neonatal circumcision: A randomized clinical trial. *JAMA* 1997;278:2157-62. <https://doi.org/10.1001/jama.1997.03550240047032>
187. Butler-O'Hara M, LeMoine C, Guillet R. Analgesia for neonatal circumcision: A randomized controlled trial of EMLA cream vs. dorsal penile nerve block. *Pediatrics* 1998;101:E5. <https://doi.org/10.1542/peds.101.4.e5>
188. Howard CR, Howard FM, Fortune K, et al. A randomized, controlled trial of a eutectic mixture of local anesthetic cream (lidocaine and prilocaine) vs. penile nerve block for pain relief during circumcision. *Am J Obstet Gynecol* 1999;181:1506-11. [https://doi.org/10.1016/S0002-9378\(99\)70397-2](https://doi.org/10.1016/S0002-9378(99)70397-2)
189. Cyna A, Middleton P. Caudal epidural block vs. other methods of postoperative pain relief for circumcision in boys. *Intervention Review; Cochrane Database Syst Rev* 2008;1:35.
190. Soh C, Ng S, Lim S. Dorsal penile nerve block. *Paediatric Anaesthesia* 2003;13:329-33. <https://doi.org/10.1046/j.1460-9592.2003.01033.x>
191. Long R, McCartan D, Cullen I, et al. A preliminary study of the sensory distribution of the penile dorsal and ventral nerves: Implications for effective penile block for circumcision. *BJU Int* 2009;105:1576-8. <https://doi.org/10.1111/j.1464-410X.2009.09044.x>
192. Stevens B, Yamada J, Ohlsson A. Sucrose for analgesia in newborn infants undergoing painful procedures (Review). *Intervention Review* 2010;1-112.
193. South M, Strauss R, South A, et al. The use of non-nutritive sucking to decrease the physiologic pain response during neonatal circumcision: A randomized controlled trial. *Am J Obs Gynecol* 2005;193:537-43. <https://doi.org/10.1016/j.ajog.2005.03.060>
194. Shah T, Raistrick J, Taylor I, et al. A circumcision service for religious reasons. *BJU Int* 1999;83:807-9. <https://doi.org/10.1046/j.1464-410x.1999.00000.x>
195. Davis DA, Mazmanian PE, Fordis M, et al. Accuracy of physician self-assessment compared with observed measures of competence: A systematic review. *JAMA* 2006;296:1094-102. <https://doi.org/10.1001/jama.296.9.1094>
196. Pieretti R, Goldstein A, Pieretti-Vanmarcke R. Late complications of newborn circumcision: A common and avoidable problem. *Pediatr Surg Int* 2010;26:515-8. <https://doi.org/10.1007/s00383-010-2566-9>
197. Kokorowski P, Routh J, Hubert K, et al. Trends in revision circumcision at pediatric hospitals. *Clin Pediatr* 2013;52:1-8. <https://doi.org/10.1177/0009922813492878>
198. Weiss H, Larke N, Halperin D, et al. Complications of circumcision in male neonates, infants and children: A systematic review. *BMC Urol* 2010;10:1-13. <https://doi.org/10.1186/1471-2490-10-2>

199. Perera CL, Bridgewater FH, Thavaneswaran P, et al. Safety and efficacy of nontherapeutic male circumcision: A systematic review. *Ann Fam Med* 2010;8:64-72. <https://doi.org/10.1370/afm.1073>
200. Neonatal circumcision revisited. Fetus and Newborn Committee, Canadian Paediatric Society. *CMAJ* 1996;154:769-80.
201. Williams N, Kapila L. Complications of circumcision. *Br J Surg* 1993;80:1231-6. <https://doi.org/10.1002/bjs.1800801005>
202. Wiswell TE, Geschke DW. Risks from circumcision during the first month of life compared with those for uncircumcised boys. *Pediatrics* 1989;83:1011-5.
203. Christakis DA, Harvey E, Zerr DM, et al. A trade-off analysis of routine newborn circumcision. *Pediatrics* 2000;105:246-9. <https://doi.org/10.1097/00006254-200009000-00010>
204. Krill A, Palmer L, Palmer J. Complications of circumcision. *Scientific World Journal* 2011;11:2458-68. <https://doi.org/10.1100/2011/373829>
205. Neonatal herpes simplex virus infection following Jewish ritual circumcisions that included direct orogenital suction — New York City, 2000-2011. *MMWR* 2012;61:405-509.
206. Joudi M, Fathi M, Hiraifar M. Incidence of asymptomatic meatal stenosis in children following neonatal circumcision. *J Pediatr Urol* 2011;7:526-8. <https://doi.org/10.1016/j.jpuro.2010.08.005>
207. Van Howe R. Incidence of meatal stenosis following neonatal circumcision in a primary care setting. *Clin Pediatr* 2006;49:54. <https://doi.org/10.1177/000992280604500108>
208. Yegane R, Kheirollahi A, Salehi N, et al. Late complications of circumcision in Iran. *Pediatr Surg Int* 2006;22:442-5. <https://doi.org/10.1007/s00383-006-1672-1>
209. Mousavi S, Salehifar E. Circumcision complications associated with the Plastibell device and conventional dissection surgery: A trial of 586 infants of ages up to 12 months. *Adv Urol* 2008;4:1-8. <https://doi.org/10.1155/2008/606123>
210. Bode C. O., Ikhisemoje S, Ademuyiwa AO. Penile injuries from proximal migration of the Plastibell circumcision ring. *J Pediatr Urol* 2010;6:23-7. <https://doi.org/10.1016/j.jpuro.2009.05.011>
211. Eroglu E, Bastian O, Ozkan H, et al. Buried penis after newborn circumcision. *J Urol* 2008;181:1841-3. <https://doi.org/10.1016/j.juro.2008.12.006>
212. Horowitz M, Gershbein AB. Gomco circumcision: When is it safe? *J Pediatr Surg* 2001;36:1047-9. <https://doi.org/10.1053/jpsu.2001.24739>
213. Storm DW, Baxter C, Koff SA, et al. The relationship between obesity and complications after neonatal circumcision. *J Urol* 2011;186:1638-41. <https://doi.org/10.1016/j.juro.2011.04.016>
214. Bastos Netto JM, de Araújo JG Jr, de Almeida Noronha MF, et al. Prospective randomized trial comparing dissection with Plastibell® circumcision. *J Pediatr Urol* 2010;6:572-7. <https://doi.org/10.1016/j.jpuro.2010.01.005>
215. Feinberg A, Brust R, Walker T. Bleeding at circumcision: Patient or operator issue? *Clin Pediatr* 2010;49:760-3. <https://doi.org/10.1177/0009922810364658>
216. Palmer J, Elder J, Palmer L. The use of betamethasone to manage the trapped penis following neonatal circumcision. *J Urol* 2005;174:1577-8. [https://doi.org/10.1016/S0022-5347\(01\)68726-3](https://doi.org/10.1016/S0022-5347(01)68726-3)
217. Blalock H. J, Vemulakonda V, Ritchey M, et al. Outpatient management of phimosis following newborn circumcision. *J Urol* 2003;169:2332-4. <https://doi.org/10.1097/01.ju.0000067602.42875.d8>
218. Griffiths DM, Atwell JD, Freeman NV. A prospective survey of the indications and morbidity of circumcision in children. *Eur Urol* 1985;11:184-7. <https://doi.org/10.1159/000472487>
219. Stenram A, Malmfors G, Okmian L. Circumcision for phimosis — indications and results. *Acta Paediatr Scand* 1986;75:321-3. <https://doi.org/10.1111/j.1651-2227.1986.tb10207.x>
220. Sansom S, Prabhu V, Hutchinson A, et al. Cost-effectiveness of newborn circumcision in reducing lifetime HIV risk among US males. *PLoS* 2010;5:1-7.
221. Schoen E, Colby C, To T. Cost analysis of neonatal circumcision in a large health maintenance organization. *J Urol* 2006;175:1111-5. [https://doi.org/10.1016/S0022-5347\(05\)00399-X](https://doi.org/10.1016/S0022-5347(05)00399-X)
222. Van Howe RS. A cost-utility analysis of neonatal circumcision. *Med Decis Making* 2004;24:584-601.
223. Kacker S, Frick K, Gaydos C, et al. Cost and effectiveness of neonatal male circumcision. *Arch Pediatr Adolesc Med* 2012;166:910-8. <https://doi.org/10.1001/archpediatrics.2012.1440>
224. Anderson J, Wilson D, Templeton D, et al. Cost-effectiveness of adult circumcision in a resource-rich setting for HIV prevention among men who have sex with men. *J Infect Dis* 2009;200:1803-12. <https://doi.org/10.1086/648472>
225. Binagwaho A, Pegurri E, Muta J, et al. Male circumcision at different ages in Rwanda: A cost-effectiveness study. *PLoS* 2010;7:1-10. <https://doi.org/10.1371/journal.pmed.1000211>
226. Kahn J, Marseille E, Auvert B. Cost-effectiveness of male circumcision for HIV prevention in a South African setting. *PLoS* 2006;3:2349-58. <https://doi.org/10.1371/journal.pmed.0030517>
227. McAllister R, Travis J, Bollinger D. The cost to circumcise Africa. *J Mens Health* 2008;7:307-16. <https://doi.org/10.3149/jmh.0703.307>
228. Roca P, Alvarado C, Stausmire J, et al. Effectiveness of a simulated training model for procedural skill demonstration in neonatal circumcision. *Simul Healthc* 2012;7:362-73. <https://doi.org/10.1097/SIH.0b013e3182634ace>
229. Freeman J, Dobbie A. Neonatal circumcision model and competency evaluation for family medicine residents. *Family Medicine* 2007;39:241-3.
230. Le B, Mickelson J, Gossett D, et al. Residency training in neonatal circumcision: A pilot study and needs assessment. *J Urol* 2010;184:1754-7. <https://doi.org/10.1016/j.juro.2010.03.077>
231. Kiggundu V, Watya S, Kigorzi G et al. The number of procedures required to achieve optimal competency with male circumcision: Findings from a randomized trial in Rakai, Uganda. *BJU Int* 2009;104:529-32. <https://doi.org/10.1111/j.1464-410X.2009.08420.x>
232. Demaria J, Abdulla A, Pemberton J, et al. Are physicians performing neonatal circumcisions well-trained? *Can Urol Assoc J* 2013;7:26-4. <https://doi.org/10.5489/cuaj.200>
233. Guyatt GH, Oxman AD, Vist GE, et al. GRADE: An emerging consensus on rating quality of evidence and strength of recommendations. *BMJ* 2008;336:924-6. <https://doi.org/10.1136/bmj.39489.470347.AD>

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