Managing the adverse events of intravesical bacillus Calmette–Guérin therapy

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Abstract: This paper provides recommendations on the management of complications arising from intravesical treatment with bacillus Calmette–Guérin (BCG) for nonmuscle-invasive bladder tumors. There is minimal recommendations currently available as randomized trials on the side effects of intravesical BCG are lacking and severe complications are usually described in case reports only. All physicians giving intravesical BCG should be aware of the possible complications that could arise and how to treat these. The incidence of bladder irritation, general malaise, and fever is very high, while severe complications remain rare. Approximately 8% of patients have to stop treatment because of these complications. BCG infections and reactions can occur anywhere in the body, and may happen straight away or even several months or years after BCG treatment, making early diagnosis difficult. Additionally, correct diagnosis is hampered by the uncertain appearance of BCG in tissue and body fluid. An essential step in the management complications arising from BCG is written information for both the family doctor and the patient on the possible adverse events and their management. Recent data demonstrated that none of the earlier advocated methods to prevent BCG toxicity are valid: lowering the dose, tuberculostatic drugs, or oxybutynin. Severe complications are treated with three or four tuberculostatics over 3–12 months, depending on the severity of the situation. Corticosteroids are an essential therapy in BCG septicemia. Nonsteroidal anti-inflammatory drugs and corticosteroids can manage efficiently the immunological complications.

Keywords: BCG, intravesical therapy, complications

Introduction
Bacillus Calmette-Guérin (BCG) has been used to treat nonmuscle-invasive bladder tumors for >40 years. It is one of the most successful biotherapies for cancer in use. Despite this long clinical experience, the mechanism of its therapeutic effect is still under investigation. Available evidence suggest that urothelial cells, including the bladder cancer cells themselves, cells of the immune system, and their secretions both have crucial roles in its antitumor effect. Several of the side effects are related to this strong immune response and others to the fact that living bacteria are instillated in the bladder.

BCG is found to be the most effective prophylactic treatment for patients with high-risk nonmuscle-invasive urothelial neoplasms, and its use is therefore recommended in international guidelines. In intermediate risk disease, it was also found more effective than intravesical chemotherapy but its use should be outweighed against its potential severe toxicity. BCG has no place in the treatment of low-grade urothelial neoplasms except in the rare cases where chemotherapy is unable to reduce the recurrence rate of the tumors. The severe side effects are a major reason to avoid
the use of BCG. Nevertheless, BCG remains an important and common treatment to avoid the need for cystectomy in urothelial cancer. Every physician treating this disease should therefore know how to prevent and manage the complications of BCG treatment.

**Previous reviews on management of BCG complications**

A committee of international experts in bladder cancer management, known as the International Bladder Cancer Group, published practical recommendations for prevention and treatment of intravesical BCG adverse events in October 2008. This was based on a review of the literature and to a large extent on expert opinion because literature did not deliver a level of evidence greater than level 3. As several of the authors of this article were also involved in previous reviews, this document reflects the opinions that were already published before 2008. Some of their recommendations did not survive. After 2008, prevention of adverse events with ofloxacin, lowering BCG dose, and use of oxybutynin were not confirmed to be effective in randomized trials (see below).

To our knowledge and after a PubMed search from January 2000 to May 2015 with the terms “BCG complications”, the current article is the last complete review with treatment recommendations published.

**Frequency and timing of the side effects**

The largest and most recent published study with BCG, including 1,316 patients, is the EORTC study comparing one-third dose with full dose BCG and 1 year with 3 years of maintenance BCG. It reports the side effects in a standard way, and also a recommendation on the treatment of the complication was provided in the study protocol. Its publication on the side effects contains interesting and recent information on BCG toxicity. Among the 1,316 patients who started BCG, 62.8% reported local side effects and 30.6% had some form of systemic side effects, from a frequent short period of fever and general malaise to rare severe systemic complications. This is in line with the previous reports on this subject and illustrates the high frequency of the side effects.

The most frequent local side effects in the EORTC study are complaints of BCG-induced cystitis (35%) and bacterial infection (23.3%), frequency of more than once per hour in 23.6%, and macroscopic hematuria in 22.6%. The most frequent systemic side effects were general malaise in 15.5% and fever in 8.1%. BCG sepsis was observed in four patients (0.3%). In contrast to the general belief that side effects increase over time, frequency was similar in the induction course (first six instillations), during the first year, and in the 2 following years. Most treatment discontinuations for severe side effects occurred in the first year (6.2% on a total of 7.8% discontinuations over the whole treatment period). Severe reactions can already appear at the first instillation, but obviously the side effects are not dependent on the number of instillations but upon the host. These results confirm the observation of a previous EORTC study (30911) where the majority of the side effects occurred within the first year, and still 19% of the patients stopped treatment because of toxicity. Obviously, the discontinuation rate diminished with increasing experience among urologists with BCG as only 7.8% stopped treatment in the last study.

**Type of severe side effects**

Severe side effects continue to be reported in the literature. BCG infection or immune reactions can appear wherever in the body, in any organ, with very usual clinical manifestations in different medical specialties. A review on the subject was published in August 2014 and gives an extensive list of described severe complications: prostatitis, orchiepididymitis, balanitis, osteomyelitis, and acute infectious arthritis at different places in the skeleton, infected orthopedic and vascular prostheses, rheumatological complications, mycobacterial pneumonia, interstitial pneumonitis due to hypersensitivity, hepatitis, nephritis, vasculitis, mycotic aneurisms, disseminated BCG and septicemia, and finally the local complication of ureteral obstruction and bladder contracture. From June 2014 to May 2015, we found another ten reports on severe complications. So the things that can happen are very rare but quite impressive and in any organ of the body.

**Difficulties in diagnosis of the severe side effects**

If the side effects occur shortly after the instillation, the relationship with BCG can be made easily. Nevertheless, the side effects mostly occur when the patient is already back at home and generalists are not familiar with the complications of BCG. Therefore, it is highly advocated to inform the family doctor of the patient on the treatment and its possible complications, alarm symptoms, and practical attitude to solve the problem.

A second problem is that complications occur late, up to years after the treatment, when the urologist, who gives it, is no longer involved. Additionally, many of the complications...
give symptoms elsewhere in the body, and the patient goes 
to specialists who, again, are unfamiliar with BCG mani-
festations or even unaware of this treatment. Therefore, an 
information document should be delivered to each patient 
treated with the possible late complications. This is the way 
we handle it since many years. Although we have no proof, 
we are convinced that this can help to come to a more rapid 
diagnosis by colleagues of other specialties. This is important 
as the outcome of a complication is often dependent on early 
treatment initiation.

Even if a BCG complication is suspected, it may be 
difficult to prove it. Acid-fast staining, culture, and pol-y-
merase chain reaction testing are often negative. Tissue 
biopsies and cultures should be performed to evaluate 
noncaseating granuloma formation and the presence of 
Mycobacterium bovis, realizing, however, that this has a 
low yield. As a result, a high clinical suspicion is critical in 
order to prevent delays in treatment initiation.

**Differences in adverse events among different BCG strains**

Side effects seem similar for all BCG strains used (Tice, 
Connaught, Pasteur, RIVM), but frequency differs from 
one study to another. No differences between strains in side 
effects have been noted in a meta-analysis and in a recent 
study comparing Tice versus Connaught strain. The dose 
and treatment schedules used differ among studies, and 
therefore, comparison among them is difficult. Ideally, the 
dose should be expressed on colony-forming units as there 
are strong variations of factors 1–4 among the commercial 
preparations. Many studies do not mention this information 
and express the dose in milligrams.

Another problem is that the side effects of BCG have 
not been evaluated in a standard way in many studies, 
resulting in large differences in the frequency of local side 
effects of BCG. A classification of these side effects, taking 
into account severity and duration, has been proposed 
and validated, but this was not commonly applied by the 
urologic community.

**Prevention strategies to diminish toxicity**

Avoid factors favoring side effects

Several of the severe general side effects could be corre-
lated with traumatic instillation or with instillations given 
early after the transurethral resection. Therefore, it is 
recommended not to start the instillations within the first 
14 days after the resection or when there is still macroscopic 
hematuria, suggesting insufficient healing of the resection 
wound. BCG should not be given after a traumatic cath-
eterization as there is an open wound that can give direct 
access to the blood circulation provoking disseminated 
BCG infection. Many experts suggest delaying treatment 
in the presence of bacterial cystitis because the barrier for 
the BCG to reach the bloodstream is traumatized. However 
severe side effects continue to appear even after respecting 
the aforementioned rules.

**Preventive systemic administration of tuberculostatic drugs**

As BCG strains are sensitive for most tuberculostatic drugs, it 
was logic to explore if its use could diminish the side effects, 
with however possible less efficacy. EORTC study (30911) 
addressed this clinical question. Three days of 300 mg iso-
niazid (or isonicotinylhydrazide [INH]), once daily at the 
occasion of the BCG instillation, could not reduce local or 
systemic toxicity and did not influence efficacy on recur-
rence and progression. However, INH provoked transient 
cholangitis in several patients. So, the use of prophylactic 
INH is not recommended.

Another prospective, double-blind, placebo-controlled 
multicenter randomized clinical trial showed that two doses 
of 200 mg ofloxacin, which is a strong tuberculostatic agent, 
given shortly after BCG instillation, reduced moderate-to-
severe side effects by 18.5%. Compliance to the therapy was 
also better. However, it concerned only 115 patients, divided 
in two groups, which makes definitive conclusions dangerous.

As the median follow-up was only 1 year, long-term efficacy 
cannot be judged. To our knowledge, further reports on this 
method of prevention did not appear so far, and therefore, it 
cannot (yet) be advocated.

**Preventive symptomatic treatment of bladder irritation during BCG**

As frequency is a major complaint during BCG instillations, 
10 mg oxybutynin extended release, twice a day, has 
been tested in a randomized study versus placebo includ-
ing 50 patients. The results were disappointing as the 
significantly worse outcome was in the oxybutynin arm.

**Lowering the BCG dose**

Low-dose BCG has been tried in an attempt to decrease the 
frequency and severity of the side effects. In this study, 
one-third dose BCG was found as effective as a full dose 
in the prevention of recurrence and progression. However, 
patients with multifocal tumors fared better with the standard
dose. In a later study, they confirmed similar efficacy of one-third dose versus full dose in high-grade and high-risk tumors, while overall side effects were significantly less. However, the number of patients who discontinued BCG for toxicity and the severe complications were similar in both arms. A third report of the same group found that one-sixth dose was significantly less effective than one-third dose, and therefore should not be used. The EORTC showed in a Phase II marker lesion study that a quarter dose was still effective. So, this is probably the lowest dose that can be advocated regarding efficacy.

The largest and most recent EORTC study confirms the data on efficacy of the one-third dose BCG but cannot confirm any difference in adverse events with the full dose. The explanation for these differences in outcome between the Spanish and EORTC study is unclear. A major difference in the study design was the duration of BCG treatment, which was 5 months in the Spanish study and 1–3 years in the EORTC study. Anyhow, it is far from evident that a lower dose will diminish side effects in general and certainly not the severe and most troublesome ones, which are the reason for stopping the treatment with BCG.

**Use of inactivated BCG**

The idea to inactivate BCG by heat, irradiation, or formaldehyde in order to prevent its side effects due to the living bacteria is already old, but recently was addressed again in preclinical studies. However, one is far from clinical application. Anyhow, one may expect that, if inactivated BCG keeps its therapeutic effect, the side effects due to the immune response will remain.

**Treatment of the BCG complications**

**BCG cystitis**

Pollakisuria, dysuria, urgency, and hematuria are frequent in the first 2 days after BCG instillation and do not need therapy. Increased diuresis, to enhance the evacuation of the mycobacteria, is advocated without scientific proof of efficacy, but it seems logic. When side effects remain for longer time or are really intolerable for the patient, symptomatic treatment with spasmolytics, anticholinergics, antiphlogistics, and analgesics are empirically advised. Except for oxybutynin, which failed to be effective, none of these drugs have been tested in this condition versus placebo. A randomized clinical trial with ofloxacin could diminish local side effects and therefore can be recommended in persisting severe cystitis. In view of the rather high frequency of associated bacterial cystitis, it is worthwhile to make a culture of the urine and to start thereafter blindly with ofloxacin. This is a pragmatic way of helping a patient who is suffering and claiming therapy. Therapy can be adjusted when the result of the urine culture is available. Anyhow, next instillation should be postponed with at least a week. There is little literature on BCG efficacy altered by changing the intensity of the therapy, but postponing the instillation is the only reasonable solution to keep the patient on further BCG treatment. All these recommendations are based on expert opinion, taken from the previous reviews, as randomized clinical trials or even large observational studies on efficacy of the recommended therapies are missing.

**General malaise and fever**

Again, these frequent side effects resolve mostly within 48 hours. Symptomatic antipyretics can be given when fever exceeds 38°C. Fever >38.5°C for >2 days needs close monitoring of the patient and consultation of infectious disease specialist in order to see if there are other causes of fever. While further diagnostic evaluation, prompt treatment with a minimum of two or more tuberculostatic agents (eg, fluoroquinolones, INH, rifampicin) is started. The duration of this therapy is badly defined and depends on further evolution of fever, malaise, and other findings at diagnostic exploration. Anyhow, further BCG is at least postponed, but with a high-risk tumor, the side effects should be weighed against the benefits.

**Local infections with BCG**

BCG can invade the prostate and the seminal vesicles up to the epididymis. These diseases require a triple tuberculostatic treatment. Often fluoroquinolone is one arm of it. The duration varies from one study to another from 3 to 6 months. Orchiepididymectomy may be a rapid solution for a severely symptomatic patient, but it does not replace the general tuberculostatic treatment that should be continued. In rare cases, transurethral resection of the prostate is necessary to solve obstruction. Asymptomatic granulomatous prostatitis, however, was found to be frequent after intravesical BCG therapy but does not require treatment. BCG balanitis or contact dermatitis has been described and can be handled with local steroids. Careful cleaning of hands and genital region after drug handling and voiding should prevent it.

Granulomatous balanitis, however, can present with multiple erythematous and painless nodules of the glans, which should be treated with systemic triple tuberculostatic drug therapy.
Systemic BCG infections
As described earlier, BCG can spread from the bladder in any organ in the body and provoke a wide range of manifestations, shortly or several months after BCG instillation. All these conditions require cessation of BCG, if still applicable, and a systemic treatment with at least three tuberculostatic agents. The duration of the treatment reported in the literature is variable from 3 months to 1 year, depending on the severity of the complication treated. This is based rather on expert opinion than on any evidence. In general, a 6-month therapy is advocated.

In case of septicemia and multiorgan failure, early high-dose corticosteroids support is an essential part of the treatment as long as symptoms persist. In these cases, it is also recommended to start as many as four tuberculostatic agents in order to obtain rapid and trustable response. Inadvertent intravenous BCG administration was also reported. This is due to the dubious instruction from the urologist who wrote “IV BCG” (meaning intravesical) for an unexperienced nurse.

Tuberculostatic agents
The tuberculostatic drugs that can be used against BCG are INH, rifampicin, ethambutol, fluoroquinolones, clarithromycin, aminoglycosides, and doxycycline. BCG strains are not sensitive to pyrazinamide, and therefore should not be used, although mentioned in several articles.

The use of cycloserine is controversial. Although it has some survival benefits in mice experiments, the drug seems to be inactive against the current strains of BCG. Its use cannot be recommended.

Monotherapy is never indicated as resistance to one drug rapidly appears. Triple drugs, and even four drugs in the severe cases, should be used. The drugs are given as a single daily dose, all together before breakfast, on an empty stomach.

Tolerability and side effects may be a serious problem. INH has a well-known liver toxicity, which appears in 10%–20% of the patients. When liver transaminases exceed three to five times the normal value, the drug must be stopped. It is also recommendable to control the transaminases before start of the treatment. Rifampicin can add to the liver toxicity and has many interactions with other drugs. Ethambutol can provoke optical neuritis. When these drugs are prescribed by physicians who are not familiar with those drugs, it is advocated to consult the professional websites on that subject or to refer patients to colleagues or centers familiar with the treatment of tuberculosis.

Immunological complications
BCG incites a strong immune reaction, and therefore, immunologically induced diseases and symptoms may be expected. More than 89 cases of arthritis and arthralgia provoked by BCG have been described in a recent review article, and a more recent case can be added. Polyarthritis is the most frequent form of the disease, but one or few articulations can be attacked. Mild and short arthralgias, without consequences, occurred regularly during a BCG course. Reiter’s syndrome and conjunctivitis alone have been described in association with the arthritis. Immunological reactions at any other organ in the body theoretically can be expected and are rarely described. The diagnostic workup of these patients may be exhaustive given the broad range of causes of similar symptoms.

Nonsteroidal anti-inflammatory drugs and corticosteroids are effective in the large majority of the cases with recovery within 2 and 6 months, respectively, of 70% and 93% of the patients.

Guidelines on nonmuscle-invasive bladder cancer
BCG is a recommended treatment in all the guidelines on nonmuscle-invasive bladder cancer. In view of the high number of side effects and a small number but severe, potentially lethal complications, it seems reasonable that they give instructions on the management of the complications of BCG. This review draws attention to the fact that some of the suggested methods to prevent BCG toxicity are no longer valid.

An important suggestion is to give written information to patients and their general practitioner on the possible severe, early, and late complications in any organ of the body. Although it has not been proved, it seems a method of common sense to improve early detection and therapy.

Conclusion
- All physicians giving intravesical BCG should be aware of the possible complications and its treatments.
- The side effects occur from the first instillation to the last one and the incidence is not increasing over time.
- The incidence of bladder irritation, general malaise, and fever is very high, while severe complications remain rare. Following the most recent study, ~8% of the patients have to stop treatment for these complications.
- BCG infections and reactions can occur at any organ or place in the body, and this can happen shortly or months to years after the BCG treatment, making early diagnosis difficult.
- Additionally, correct diagnosis is hampered by uncertain appearance of BCG in tissues and body fluids.
An essential step in the management of BCG complications is written information for the family doctor and for the patient on the possible adverse events and their management.

Recent data demonstrate that none of the earlier suggested methods to prevent BCG toxicity are valid: lowering the dose, tuberculostatic drugs, or oxybutynin.

The most common side effects do not need therapy.

Severe complications are treated with three or four tuberculostatic drugs during 3–12 months depending on the severity of the situation.

Corticosteroids are an essential therapy in BCG septicemia. Nonsteroidal anti-inflammatory drugs and corticosteroids resolve the immunological complications, mostly rheumatological, of BCG.

**Disclosure**

The authors report no conflicts of interest in this work.

**References**


