Reduced Dose Intravesical Bacillus Calmette-Guérin: Why It Might Not Matter

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Abstract. When it comes to the treatment of patients with non-muscle-invasive bladder cancer (NMIBC) with intravesical bacillus Calmette-Guérin (BCG), two questions must be considered: 1) what dose to give, and 2) for how long? The issue of optimal dose and duration has been the subject of several randomized trials and is especially pertinent in the context of a global BCG shortage. Despite this, there appears to be uncertainty as to whether BCG dose or duration may be compromised in the event of shortage. As such, we wish to summarize the available evidence as an aid to the practicing urologist.

Keywords: Non-muscle-invasive bladder cancer, bacillus Calmette-Guerin, reduced dose, immunotherapy

BACKGROUND

One of the earliest randomized trials to address the question of BCG dose was conducted by the CUETO group who compared full dose BCG (81mg) with one-third dose (27mg) in intermediate- and high-risk patients with NMIBC [1]. Both dose cohorts received a total of 12 instillations – a 6-week induc-
tion course followed by a further six instillations given fortnightly – and found similar recurrence rates (29.4% in the full dose arm vs. 30.7% in the reduced dose arm) with no significant difference in time to first recurrence \( (p = 0.586) \). Similarly, both cohorts had similar rates of progression to muscle-invasive disease (11.5% vs. 13.3%). When looking for clues as to subgroup analysis, they observed a trend towards increased efficacy with full dose BCG against progression in patients with multifocal disease \( (p = 0.048) \). In a later study, the CUETO group addressed the same question in patients with high-risk NMIBC (G3/T1/Tis), and once again demonstrated that one-third dose BCG was as effective as full dose [2]. Subsequently, the EORTC 30962 study examined the use of full- and one-third dose OncoTICE® BCG given as either a 1- or 3-year maintenance course in 1355 patients with intermediate- and high-risk NMIBC [3]. At a median follow-up of 7.1 years, the authors reported that full dose BCG was not superior to one-third dose BCG (5-year disease-free rate: 61.7% vs. 58.5%, \( p = 0.092 \)). When looking at subgroups, in high-risk patients, full dose BCG given for 3-years reduced recurrences compared with full dose for 1-year; however, there were no differences in progression or survival. Additionally, a recent retrospective analysis of 563 patients with intermediate- and high-risk NMIBC treated with either reduced or full dose adequate BCG during the BCG shortage provided

real-world experience that the use of one-third dose BCG was not associated with adverse oncological outcomes [4]: time to recurrence \( (p = 0.449) \), time to progression (0.716) and cancer-specific survival \( (p = 0.320) \) was similar in both groups.

The results of these clinical trials are not unexpected. Each trial evaluated full and reduced dose BCG with regards to the clinically used supply, which is by weight of the lyophilized powder. What is not widely known is that the number of viable BCG organisms contained in clinical supply BCG is expressed as a range with an almost ten fold variation in amount of organisms in each vial. For example, using the data from TICE® BCG (Merck, USA), the main strain in North America and many other parts of the world, we know that one vial of TICE® BCG contains between 1 to \( 8 \times 10^8 \) colony forming units (CFU) [4]. Put in other terms, each milligram of lyophilized BCG contains 2 million to 16 million CFUs of actual BCG organisms. Given this wide variation in CFUs, using one-third dose will not necessarily result in suboptimal dosing.

In a hypothetical situation, patient X receives full dose BCG from a vial containing 2 million CFUs of BCG per milligram. Thus s/he received \( 50 \times 2 = 100 \) million CFUs per instillation (Fig. 1). Patient Y on the other hand, receives ‘reduced dose’ BCG from a vial containing 16 million CFUs of BCG. Thus s/he received \( 16.6 \times 16 = 256 \) million CFUs per instillation, which is greater than the ‘full dose’ received.

![Dose delivered (million CFUs/instillation)](image)

Fig. 1. Patient X receives “full dose” (BCG 50 mg) from a vial containing 2 million CFUs/mg, thus receiving 100 CFUs per instillation. Patient Y receives “reduced-dose” BCG (16.6 mg) from a vial containing 16 million CFUs/mg, thus receiving 256 million CFUs per instillation. In this situation, the patient receiving 1/3rd dose actually is getting 2.5 times the dose of the ‘full dose’ patient.
by patient X. Of course, each manufacturer of BCG might have their own range of CFU variation (e.g., Onco-BCG® from the Serum Institute of India has an even greater variation of $1 \sim 19.2 \times 10^8$ CFUs per vial) so we urge readers to read up on the specifics of this variation for the strain of BCG used in their centers (e.g., from manufacturer data sheets).

**IMPLICATIONS**

It is important to clarify that this letter serves to provide an explanation as to why patients treated with reduced dose BCG may not actually receive fewer CFUs than patients receiving full dose BCG. With this in mind, what are the implications for our patients? Firstly, patients may be reassured that if they do receive reduced dose BCG, based on data from RCTs, this likely has no major impact on their disease-related outcomes. Indeed, recent American Urological Association/Society of Urologic Oncology guidelines for the treatment of NMIBC recommend the use of reduced dose BCG for maintenance treatment [5]. Secondly, during these times of BCG shortage, splitting a vial of BCG amongst three patients allows more patients to receive BCG. Since the preponderance of data suggests that receipt of maintenance BCG is an important component of the durability of response — especially with regards to decreasing progression rates — it is preferable that patients receive a reduced dose (either one-half or one-third dose) with the standard 6 + 3 schedule rather than full dose BCG given over a shorter duration. We acknowledge that some institutions may be hesitant to employ split-vial dosing given that manufacturers stipulate BCG should be administered within 2 hours of reconstitution. However, it has been shown that BCG remains viable for at least 8 hours and, in some cases, up to 72 hours after reconstitution [6]. Furthermore, whilst the 3 weekly maintenance schedule is important, this timing is an approximation rather than a rigid prescription. As such, providers have some flexibility around the scheduling of maintenance treatments and may split the dose based on when maximal patients are available. With that said, one-third dose appears to be the minimum required for clinical effectiveness as demonstrated by the CUETO group who showed that one-sixth dose BCG was inferior to one-third dose [7]. Finally, this carries practical implications for the conduct of clinical trials for BCG-unresponsive disease. While the United States Food and Drug Administration does not specify that patients must receive full dose BCG in order to qualify for trials of agents in the BCG-unresponsive setting, many sponsors are hesitant to include patients who have received one-third dose BCG for induction and/or maintenance during these times of shortage. We believe, as previously stated [8], that these patients should be allowed to enroll in trials for the aforementioned reasons.

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REFERENCES


