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Efficient Random-Walk Methods for Approximating Polytope Volume

Ioannis Z. Emiris*

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Abstract

We experimentally study the fundamental problem of computing the volume of a convex polytope given as an intersection of linear inequalities. We implement and evaluate practical randomized algorithms for accurately approximating the polytope's volume in high dimensions (e.g. one hundred). To carry out this efficiently we experimentally correlate the effect of parameters, such as random walk length and number of sample points, on accuracy and runtime. Moreover, we exploit the problem's geometry by implementing an iterative rounding procedure, computing partial generations of random points and designing fast polytope boundary oracles. Our publicly available code is significantly faster than exact computation and more accurate than existing approximation methods. We provide volume approximations for the Birkhoff polytopes $\mathcal{B}_{11}, \dots, \mathcal{B}_{15}$, whereas exact methods have only computed that of \mathcal{B}_{10} .

Keywords: volume approximation, general dimension, random walk, polytope oracle, algorithm engineering, ray shooting

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1 Introduction

A fundamental problem in discrete and computational geometry is to compute the volume of a convex body in general dimension or, more particularly, of a polytope. In the past 15 years, randomized algorithms for this problem have witnessed a remarkable progress. Starting with the breakthrough poly-time algorithm of [15], subsequent results brought down the exponent on the dimension from 27 to 4 [28]. However, the question of an efficient implementation had remained open.

Notation. Convex bodies are typically given by a membership oracle. A polytope $P \subseteq \mathbb{R}^d$ can also be represented as the convex hull of vertices (V-polytope) or, as is the case here, as the (bounded) intersection

$$P := \{x \in \mathbb{R}^d \mid Ax \leq b\}$$

of m halfspaces given by $A \in \mathbb{R}^{m \times d}$, $b \in \mathbb{R}^m$ (H-polytope); ∂P is its boundary, and $O^*(\cdot)$ hides polylog factors in the argument. The input includes approximation factor $\epsilon > 0$; W denotes the most important runtime parameter, namely random walk length.

Previous work. Volume computation is $\#\text{-P}$ hard for V- and for H-polytopes [16]. Several exact algorithms are surveyed in [9] and implemented in VINCI [8], which however cannot handle general polytopes for dimension $d > 15$. An interesting challenge is the volume of the n -Birkhoff polytope, computed only for $n \leq 10$ using highly specialized software (Sect. 4). Regarding deterministic approximation, no poly-time algorithm can compute the volume with less than exponential relative error [17]. The algorithm of [7] has error $\leq d!$.

The landmark randomized poly-time algorithm in [15] approximates the volume of a convex body with high probability and arbitrarily small relative error. The best complexity, as a function of d , given a membership oracle, is $O^*(d^4)$ oracle calls [28]. All approaches except [28] define a sequence of co-centric balls, and produce uniform point samples in their intersections with P to approximate the volume of P .

Concerning existing software (cf Sect. 5), [13] presented recently `Matlab` code based on [28] and [12]. The latter offers a randomized algorithm for Gaussian volume (which has no direct reduction to or from volume) in $O^*(d^3)$, as a function of d . In [26] they implement [28], focusing on variance-decreasing techniques, and an empirical estimation of mixing time. In [25], they use a straightforward acceptance-rejection method, which is not expected to work in high dimension; it was tested only for $d \leq 4$. An approach using thermodynamic integration [21] offers only experimental guarantees on runtime and accuracy.

The key ingredient of all approaches is random walks that produce an almost uniform point sample. Such samples is a fundamental problem of independent interest with important applications in, e.g., global optimization, statistics, machine learning, Monte Carlo (MC) integration, and non-redundant constraint identification. Several questions of sampling combinatorial structures such as contingency tables and more generally lattice points in polytopes may be reduced to sampling a polytope.

No simple sampling method exists unless the body has standard shape, e.g., simplex, cube or ellipsoid. Acceptance-rejection techniques are inefficient in high dimensions. E.g., the number of uniform points one needs to generate in a bounding box before finding one in P is exponential in d . A Markov chain is the only known method, and it may use geometric random walks such as the grid walk, the ball walk (or variants such as the Dikin walk), and Hit-and-run [34]. The Markov chain has to make a (large) number of steps, before the generated point becomes distributed approximately uniformly (which is the stationary limit distribution of the chain). We focus on Hit-and-run which yields the fastest algorithms today.

In contrast to other walks, Hit-and-run is implemented by computing the intersection of a line with ∂P . In general, this reduces to binary search on the line, calling membership at every step. For H-polytopes, the intersection is obtained by a *boundary oracle*; for this, we employ ray-shooting with respect to the m facet hyperplanes (Sect. 2). In exact form, it is possible to

avoid linear-time queries by using space in $o(m^{\lfloor d/2 \rfloor})$, achieving queries in $O(\log m)$ [33]. Duality reduces oracles to (approximate) ε -nearest neighbor queries, which take $O(dm^{(1+\varepsilon)^{-2}+o(1)})$ using $O(dm + m^{1+(1+\varepsilon)^{-2}+o(1)})$ space by locality sensitive hashing [2]. Moreover, space-time tradeoffs from $O(1/\varepsilon^{(d-O(1))/8})$ time and $O(1/\varepsilon^{(d-O(1))/2})$ space to $O(1)$ time and $O(1/\varepsilon^{(d-O(1))})$ space are available by [3]. Approximate oracles are also connected to polytope approximation. Classic results, such as Dudley’s, show that $O((1/\varepsilon)^{(d-1)/2})$ facets suffice to approximate a convex body of unit diameter within a Hausdorff distance of ε . This is optimized to $O(\sqrt{\text{vol}(\partial P)}/\varepsilon^{(d-1)/2})$ [3]. The boundary oracle is dual to finding the extreme point in a given direction among a known pointset. This is ε -approximated through ε -coresets for measuring extent, in particular (directional) width, but requires a subset of $O((1/\varepsilon)^{(d-1)/2})$ points [1]. The exponential dependence on d or the linear dependence on m make all aforementioned methods of little practical use. Ray shooting has been studied in practice only in low dimensions, e.g., in 6-dimensional V-polytopes [37].

Contribution. We implement and experimentally study efficient algorithms for approximating the volume of polytopes. Point sampling, which is the bottleneck of these algorithms, is key in achieving poly-time complexity and high accuracy. To this end, we study variants of Hit-and-run. It is widely believed that the theoretical bound on W is quite loose, and this is confirmed by our experiments, where we set $W = O(d)$ and obtain a $< 2\%$ error in up to 100 dimensions (Sect. 4).

Our emphasis is to exploit the underlying geometry. Our algorithm uses the recursive technique of co-centric balls (cf. Sect. 3) introduced in [15] and used in a series of papers, with the most recent to be [22]. This technique forms a sequence of *diminishing* radii which, unlike previous papers, allowing us to only sample *partial generations* of points in each intersection with P , instead of sampling N points for each. In fact, the algorithm starts with computing the largest interior ball by an LP. Unlike most theoretical approaches, that use an involved rounding procedure, we sample a set of points in P and compute the minimum enclosing ellipsoid of this set, which is then linearly transformed to a ball. This procedure is repeated until the ratio of the minimum over the maximum ellipsoid axes reaches some user-defined threshold. This *iterative rounding* allows us to handle skinny polytopes efficiently.

We study various oracles (Sect. 2). Line search using membership requires $O(md + \log \frac{r}{\varepsilon_s})$ arithmetic operations. This is improved to a boundary oracle in $O(md)$ by avoiding membership. Using Coordinate Direction Hit-and-run, we further improve the oracle to $O(m)$ amortized complexity. We also exploit duality to reduce the oracle to ε -nearest neighbor search: although the asymptotic complexity is not improved, for certain instances such as cross-polytopes in $d = 16$, kd-trees achieve a 40x speed-up.

Our C++ code is open-source (sourceforge) and uses the CGAL library. A series of experiments establishes that it handles dimensions substantially larger than existing exact approaches, e.g., cubes and products of simplices within an error of 2% for $d \leq 100$, in about 20 min. Compared to approximate approaches, it computes significantly more accurate results. It computes in few hours volume estimations within an error of 2% for Birkhoff polytopes $\mathcal{B}_2, \dots, \mathcal{B}_{10}$; $\text{vol}(\mathcal{B}_{10})$ has been exactly computed by specialized parallel software in a sequential time of years. More interestingly, it provides volume estimations for $\text{vol}(\mathcal{B}_{11}), \dots, \text{vol}(\mathcal{B}_{15})$, whose exact values are unknown, within 9 hours. In conclusion, we claim that the volume of general H-polytopes in high dimensions (e.g. one hundred) can be efficiently and accurately approximated on standard computers.

Paper organization. The next section discusses walks and oracles. Sect. 3 presents the overall volume algorithm. Sect. 4 discusses our experiments, and we conclude with open questions in Sect. 5.

2 Random walks and Oracles

This section introduces the paradigm of Hit-and-run walks and focuses on their implementation, with particular emphasis on exploiting the geometry of H-polytopes. The methods presented here are analysed experimentally in Sect. 4.

Hit-and-run random walks. The main method to randomly sample a polytope is by (geometric) random walks. We shall focus on variants of Hit-and-run, which generate a uniform distribution of points [35]. Assume we possess procedure $\text{Line}(p)$, which returns line ℓ through point $p \in P \subseteq \mathbb{R}^d$; ℓ will be specified below. The main procedure of Hit-and-run is $\text{Walk}(p, P, W)$, which reads in point $p \in P$ and repeats W times: (i) run $\text{Line}(p)$, (ii) move p to a random point uniformly distributed on $P \cap \ell$. We shall consider two variants of Hit-and-run.

In *Random Directions Hit-and-run* (RDHR), $\text{Line}(p)$ returns ℓ defined by a random vector uniformly distributed on the unit sphere centered at p . The vector coordinates are drawn from the standard normal distribution. RDHR generates a uniformly distributed point in

$$O^*(d^2r^2), \text{ or } O^*(d^3r^2) \text{ oracle calls,} \quad (1)$$

with hidden constants 10^{30} , or 10^{11} respectively,

starting at an arbitrary, or at a uniformly distributed point (also known as warm start), respectively, where r is the ratio of the radius of the smallest enclosing ball over that of the largest enclosed ball in P [27].

In *Coordinate Directions Hit-and-run* (CDHR), $\text{Line}(p)$ returns ℓ defined by a random vector uniformly distributed on the set $\{e_1, \dots, e_d\}$, where $e_i = (0, \dots, 0, 1, 0, \dots, 0)$, $i = 1, \dots, d$. This is a continuous variant of the Grid walk. As far as the authors know, the mixing time has not been analyzed. We offer experimental evidence that CDHR is faster than RDHR and sufficiently accurate. An intermediate variant is Artificially Centering Hit-and-run [23], where first a set S of sample points is generated as with RDHR, then $\text{Line}(p)$ returns ℓ through p and a randomly selected point from S . This however is not a Markov chain, unlike CDHR and RDHR.

Procedure $\text{Walk}(p, P, W)$ requires at every step an access to a *boundary oracle* which computes the intersection of line ℓ with ∂P . In the sequel we discuss various implementations of this oracle.

Boundary oracle by membership. For general convex bodies, a boundary oracle can be implemented using a *membership oracle* which, given vector $y \in \mathbb{R}^d$, decides whether $y \in P$. The intersection of ℓ with ∂P is computed by binary search on the segment defined by any point on ℓ lying in the body and the intersection of ℓ with a bounding ball. Each step calls membership to test whether the current point is internal, and stops when some accuracy ϵ_s is certified. Checking the point against a hyperplane takes $O(d)$ operations, thus obtaining the intersection of ℓ with the hyperplane. We store this intersection so that subsequent tests against this hyperplane take $O(1)$. The total complexity is $O(md + \log \frac{r}{\epsilon_s})$ arithmetic operations, where r is the ball radius.

Boundary oracle by facet intersection. Given an H-polytope P the direct method to compute the intersection of line ℓ with ∂P is to examine all m hyperplanes. Let us consider $\text{Walk}(p_0, P, W)$ and line $\ell = \{x \in \mathbb{R}^d : x = \lambda v + p_0\}$, where $p_0 \in \mathbb{R}^d$ lies on ℓ , and v is the direction of ℓ . We compute the intersection of ℓ with the i -th hyperplane $a_i x = b_i$, $a_i \in \mathbb{R}^d, b_i \in \mathbb{R}$, namely $p_i := p_0 + \frac{b_i - a_i p_0}{a_i v} v$, $i \in \{1, \dots, m\}$. We seek points p^+, p^- at which ℓ intersects ∂P , namely $p^+ v = \min_{1 \leq i \leq m} \{p_i v \mid p_i v \geq 0\}$ and $p^- v = \max_{1 \leq i \leq m} \{p_i v \mid p_i v \leq 0\}$. This is computed in $O(md)$ arithmetic operations. In practice, only the λ^\pm are computed, where $p^\pm = p_0 + \lambda^\pm v$.

In the context of the volume algorithm (Sect.3), the intersection points of ℓ with ∂P are compared to the intersections of ℓ with the current sphere. Assuming the sphere is centered at the origin with radius R , its intersections with ℓ are $p = p_0 + \lambda v$ such that $\lambda^2 + 2\lambda p_0 v + |p_0|^2 - R^2 = 0$. If λ^+, λ^- give a negative sign when substituted to the aforementioned equation then p^+, p^- are the endpoints of the segment of ℓ lying in the intersection of P and the current ball. Otherwise,

we have to compute one or two roots of the aforementioned equation since the segment has one or two endpoints on the sphere.

However, in CDHR, where ℓ and v are vertical, after the computation of the first pair p^+, p^- , all other pairs can be computed in $O(m)$ arithmetic operations. This is because two sequential points produced by the walk differ only in one coordinate. Let j, k be the walk coordinate of the previous and the current step respectively. Then, assuming $P = \{x \in \mathbb{R}^d : Ax \leq b\}$, where $A \in \mathbb{R}^{m \times d}$, $\lambda^\pm = \max\{\lambda \mid A(p_0 \pm \lambda v) \leq b\}$. This becomes $\pm \lambda A v = \pm \lambda A_j \leq -A p_0 + b$, where A_j is the j -th column of A . The two maximizations are solved in $O(md)$ ops. Let vector $t = -A p_0 + b \in \mathbb{R}^m$. At the next step, given point $p'_0 = p_0 + c e_j$, where e_j is the j -th standard basis vector, we perform two maximizations $\lambda : \pm \lambda A_k \leq t - c A_j$ in $O(m)$.

Boundary oracle by duality. Duality reduces the problem to nearest neighbor (NN) search and its variants. Given a pointset $B \subseteq \mathbb{R}^d$ and query point q , NN search returns a point $p \in B$ s.t. $\text{dist}(q, p) \leq \text{dist}(q, p')$ for all $p' \in B$, where $\text{dist}(q, p)$ is the Euclidean distance between points q, p . Let us consider, w.l.o.g., boundary intersection for line ℓ parallel to the x_d -axis: $\ell = \{x : x = \lambda v + p, \lambda \geq 0\}$, $v = (0, \dots, 0, -1)$. It reduces to two ray-shooting questions; it suffices to describe one, namely with the upward vertical ray, defined by $\lambda \leq 0$. We seek the first facet hyperplane hit which, equivalently, has the maximum negative signed vertical distance from p to any hyperplane H of the upper hull, for fixed v . This distance is denoted by $\text{sv}(p, H)$. Let us consider the standard (aka functional) duality transform between points p and non-vertical hyperplanes H :

$$\begin{aligned} p = (p_1, \dots, p_d) &\mapsto p^* : x_d = p_1 x_1 + \dots + p_{d-1} x_{d-1} - p_d, \\ H : x_d = c_1 x_1 + \dots + c_{d-1} x_{d-1} + c_0 &\mapsto H^* = (c_1, \dots, c_{d-1}, -c_0). \end{aligned}$$

This transformation is self-dual, preserves point-hyperplane incidences, and negates vertical distance, hence $\text{sv}^*(p^*, H^*) = -\text{sv}(p, H)$, where $\text{sv}^*(\cdot, \cdot)$ is the signed vertical distance from hyperplane p^* to point H^* in dual space. Hence, our problem is equivalent to minimizing $\text{sv}^*(p^*, H^*) \geq 0$. Equivalently, we seek point H^* minimizing absolute vertical distance to hyperplane p^* on its side of positive distances. In dual space, consider

$$\begin{aligned} \text{point } t &= (t_1, \dots, t_d), \text{ and hyperplane} \\ p^* = q &: x_d = q_1 x_1 + \dots + q_{d-1} x_{d-1} + q_0 : \end{aligned} \tag{2}$$

$$\begin{aligned} \text{sv}^*(q, t) &= t_d - (q_1 t_1 + \dots + q_{d-1} t_{d-1} + q_0) \\ &= -(q_0, q_1, \dots, q_{d-1}, -1) \cdot (1, t_1, \dots, t_{d-1}, t_d), \end{aligned}$$

where the latter operation is inner product in Euclidean space \mathbb{R}^{d+1} of “lifted” datapoint $t' = (1, t_1, \dots, t_{d-1}, t_d)$ with “lifted” query point $q' = (q_0, q_1, \dots, q_{d-1}, -1)$. Let

$$q'' = (q', 0), \quad t'' = (t', \sqrt{M - \|t'\|_2^2}), \quad \text{for } M \geq \max_t \{1 + \|t\|_2^2\},$$

following an idea of [4]. By the cosine rule,

$$\text{dist}_{d+2}^2(q'', t'') = \|q'\|_2^2 + M + 2\text{sv}^*(q, t),$$

where $\text{dist}_{d+2}(\cdot, \cdot)$ stands for Euclidean distance in \mathbb{R}^{d+2} . Since the t'' lie on hyperplane $x_1 = 1$, optimizing $\text{dist}_{d+2}(q'', t'')$ over a set of points t'' is equivalent to optimizing $\text{dist}_{d+1}(\hat{q}, \hat{t})$, $\hat{q} = (q_1, \dots, q_{d-1}, -1, 0)$, over points $\hat{t} = (t, \sqrt{M - 1 - \|t\|_2^2})$. Hence, point t minimizing $\text{sv}^*(q, t) \geq 0$ corresponds to \hat{t} minimizing $\text{dist}_{d+1}^2(\hat{q}, \hat{t})$. Thus the problem is reduced to (exact) nearest neighbor in \mathbb{R}^{d+1} . Ray shooting to the lower hull with same v reduces to farthest neighbor. Unfortunately, an approximate solution to these problems incurs an additive error to the corresponding original problem.

Alternatively, we shall consider hyperplane queries. Let us concentrate on hyperplanes supporting facets on the lower hull of P . Their dual points lie in convex position. Given that

point p is interior in P , the dual points of the lower hull facets lie on the upper halfspace of p^* . In dual space, consider point t and hyperplane q as in expression (2). Let $\text{sd}^*(q, t)$ be the signed Euclidean distance from q to t , i.e. the minimum Euclidean distance of any point on q to t . Then $\text{sv}^*(q, t) = \text{sd}^*(q, t) / \|(q_1, \dots, q_{d-1}, 1)\|_2$, where the normal is $(q_1, \dots, q_{d-1}, 1)$. Our question, therefore, becomes equivalent to minimizing $\text{sd}^*(q, t)$ over all datapoints $t \in \mathbb{R}^d$ for which $\text{sd}^*(q, t) \geq 0$; i.e., we seek the NN above q . Starting with facets on the upper hull, the problem becomes that of maximizing $\text{sd}^*(q, t) \leq 0$, i.e. finding the NN below q .

The above approaches motivate us to use NN software for exact point and hyperplane queries (Sect. 4).

3 The volume algorithm

This section details our poly-time methods for approximating the volume of P . Algorithms in this family are the current state-of-the-art with respect to asymptotic complexity bounds. Moreover, they can achieve any approximation ratio given by the user, i.e., they form a fully polynomial randomized approximation scheme (FPRAS). Given polytope $P \subseteq \mathbb{R}^d$, they execute *sandwiching* and *Multiphase Monte Carlo* (MMC) [34].

We consider that P is a full-dimensional H -polytope. However, we can also consider P to be lower dimensional and be given in form $\{x \in \mathbb{R}^d \mid Ax = b, x \geq 0\}$, where $A \in \mathbb{R}^{m \times d}$, $x \in \mathbb{R}^d$, $b \in \mathbb{R}^m$, $A' \in \mathbb{R}^{m \times m-d+1}$, $x' \in \mathbb{R}^{m-d+1}$. Using *Gauss-Jordan elimination* the linear system $Ax = b$ can be transformed to its unique *reduced row echelon form* $[I|A']x = b'$, where I is the identity matrix. Then P can be written as $\{x' \in \mathbb{R}^{m-d+1} \mid A'x' \geq b', x' \geq 0\}$, i.e. a full-dimensional H -polytope in \mathbb{R}^{m-d+1} .

Rounding and sandwiching. This stage involves first rounding P to reach a near isotropic position, second sandwiching, i.e. to compute ball B and scalar ρ such that $B \subseteq P \subseteq \rho B$. There is an abundance of methods in literature for rounding and sandwiching (cf. [34] and references therein). However, here we develop a simple, efficient method that succeed significantly accurate results in practice (cf. Sect. 4 and Table 5). The method doesn't compute a ball that covers P but a ball B' such that $B' \cap P$ contains almost all the volume of P .

For rounding, we sample a set S of $O(n)$ random points in P . Then we approximate the minimum volume ellipsoid \mathcal{E} that covers S , and satisfies the inclusions $\frac{1}{(1+\varepsilon)^d} \mathcal{E} \subseteq \text{conv}(S) \subseteq \mathcal{E}$, in time $O(nd^2(\varepsilon^{-1} + \ln d + \ln \ln n))$ [24]. Let us write

$$\begin{aligned} \mathcal{E} &= \{x \in \mathbb{R}^d \mid (x - c_{\mathcal{E}})^T E (x - c_{\mathcal{E}}) \leq 1\} \\ &= \{x \in \mathbb{R}^d \mid L^T (x - c_{\mathcal{E}}) \leq 1\}, \end{aligned} \tag{3}$$

where $E \subseteq \mathbb{R}^{d \times d}$ is a positive semi-definite (p.s.d.) matrix and $L^T L$ its Cholesky decomposition. By substituting $x = (L^T)^{-1} y + c_{\mathcal{E}}$ we map the ellipsoid to the ball $\{y \in \mathbb{R}^d \mid y^T y \leq 1\}$. Applying this transformation to P we have $P' = \{y \in \mathbb{R}^d \mid A(L^T)^{-1} y \leq b - A c_{\mathcal{E}}\}$ which is the rounded polytope, where $\text{vol}(P) = \det(L^T)^{-1} \text{vol}(P')$. We iterate this procedure until the ratio of the minimum over the maximum ellipsoid axes reaches some user defined threshold.

For sandwiching P we first compute the *Chebyshev ball* $B(c, r)$ of P , i.e. the largest inscribed ball in P . It suffices to solve the LP: $\{\text{maximize } R, \text{ subject to: } A_i x + R \|A_i\|_2 \leq b_i, i = 1, \dots, m, R \geq 0\}$, where A_i is the i -th row of A , and the optimal values of R and $x \in \mathbb{R}^d$ yield, respectively, the radius r and the center c of the Chebyshev ball.

Then we may compute a uniform random point in $B(c, r)$ and use it as a start to perform a random walk in P , eventually generating N random points. Now, compute the largest distance between each of the N points and c ; this defines a (approximate) bounding ball. Finally, define the sequence of balls $B(c, 2^{i/d})$, $i = \alpha, \alpha + 1, \dots, \beta$, where $\alpha = \lfloor d \log r \rfloor$ and $\beta = \lceil d \log \rho \rceil$.

Multiphase Monte Carlo (MMC). MMC constructs a sequence of bodies $P_i := P \cap B(c, 2^{i/d})$, $i = \alpha, \alpha + 1, \dots, \beta$, where $P_{\alpha} = B(c, 2^{\alpha/d}) \subseteq B(c, r)$ and P_{β} (almost) contains P .

Then it approximates $\text{vol}(P)$ by the telescopic product

$$\text{vol}(P_\alpha) \prod_{i=\alpha+1}^{\beta} \frac{\text{vol}(P_i)}{\text{vol}(P_{i-1})}, \text{ where } \text{vol}(P_\alpha) = \frac{2\pi^{d/2}(2^{\lfloor \log r \rfloor})^d}{d\Gamma(d/2)}.$$

This reduces to estimating the ratios $\text{vol}(P_i)/\text{vol}(P_{i-1})$, which is achieved by generating N uniformly distributed points in P_i and by counting how many of them fall in P_{i-1} .

For point generation we use random walks as in Sect. 2. We set the walk length $W = \lfloor 10 + d/10 \rfloor = O(d)$, which is of the same order as in [26] but significantly lower than theoretical bounds. This choice is corroborated experimentally (Sect. 4).

Unlike typical approaches, which generate points in P_i for $i = \alpha, \alpha+1, \dots, \beta$, here we proceed inversely. First, let us describe initialization. We generate an (almost) uniformly distributed random point $p \in P_\alpha$, which is easy since $P_\alpha = B(c, 2^{\alpha/d}) \subseteq B(c, r)$. Then we use p to start a random walk in $P_\alpha, P_{\alpha+1}, P_{\alpha+2}$ and so on, until we obtain a uniformly distributed point in P_β . We perform N random walks starting from this point to generate N (almost) uniformly distributed points in P_β and then count how many of them fall into $P_{\beta-1}$. This yields an estimate of $\text{vol}(P_\beta)/\text{vol}(P_{\beta-1})$. Next we keep the points that lie in $P_{\beta-1}$, and use them to start walks so as to gather a total of N (almost) uniformly distributed points in $P_{\beta-1}$. We repeat until we compute the last ratio $\text{vol}(P_{\alpha+1})/\text{vol}(P_\alpha)$.

The implementation is based on a data structure S that stores the random points. In step $i > \alpha$, we wish to compute $\text{vol}(P_{\beta-i})/\text{vol}(P_{\beta-i-1})$ and S contains N random points in $P_{\beta-i+1}$ from the previous step. The computation in this step consists in removing from S the points not in $P_{\beta-i}$, then sampling $N - \text{size}(S)$ new points in $P_{\beta-i}$ and, finally, counting how many lie in $P_{\beta-i-1}$. Testing whether such a point lies in some P_i reduces to testing whether $p \in B(2^{i/d})$ because $p \in P$.

One main advantage of our method is that it creates partial generations of random points for every new body P_i , as opposed to having always to generate N points. This has a significant effect on runtime since it reduces it by a constant raised to β . Partial generations of points have been used in convex optimization [6].

We use *threads*, also in [26], to ensure independence of the points. A thread is a sequence of points each generated from the previous point in the sequence by a random walk. The first point in the sequence is uniformly distributed in the ball inscribed in P . Alg. 1 describes our algorithm using a single thread.

Complexity. The first $O^*(d^5)$ algorithm was in [22], using a sequence of subsets defined as the intersection of the given body with a ball. It uses isotropic sandwiching to bound the number of balls by $O^*(d)$, it samples $N = 400\epsilon^{-2}d \log d = O^*(d)$ points per ball, and follows a ball walk to generate each point in $O^*(d^3)$ oracle calls. Interestingly, both sandwiching and MMC each require $O^*(d^5)$ oracle calls. Later the same complexity was obtained by Hit-and-run under the assumption the convex body is well sandwiched.

Proposition 1. [22] *Assuming $B(0, 1) \subseteq P \subseteq B(0, \rho)$, the volume algorithm of [22] returns an estimation of $\text{vol}(P)$, which lies between $(1 - \epsilon)\text{vol}(P)$ and $(1 + \epsilon)\text{vol}(P)$, with probability $\geq 3/4$, by*

$$O\left(\frac{d^4 \rho^2}{\epsilon^2} \ln d \ln \rho \ln^2 \frac{d}{\epsilon}\right) = O^*(d^4 \rho^2)$$

oracle calls with probability $\geq 9/10$, where we have assumed ϵ is fixed. Sandwiching yields $\rho = \sqrt{d/\log(1/\epsilon)}$, implying a total of $O^(d^5)$ calls.*

In [28], they construct a sequence of log-concave functions and estimate ratios of integrals, instead of ratios of balls, using simulated annealing. The complexity reduces to $O^*(d^4)$ by decreasing both number of phases and number of samples per phase to $O^*(\sqrt{d})$. Using Hit-and-run, $O^*(d^3)$ still bounds the time to sample each point. Moreover, they improve isoperimetric sandwiching to $O^*(d^4)$.

Algorithm 1: VolEsti (P, ϵ, t_r)

Input : H-polytope P , objective approximation ϵ , rounding threshold t_r
Output: approximation of $\text{vol}(P)$

$N \leftarrow 400\epsilon^{-2}d \log d$; $W \leftarrow \lfloor 10 + d/10 \rfloor$;

// rounding and sandwiching
compute the Chebychev ball $B(c, r)$;
generate a random point p in $B(c, r)$;
repeat
 $S \leftarrow \emptyset$;
 for $i = 1$ to N **do**
 $p \leftarrow \text{Walk}(p, P, W)$;
 add p in S ;
 compute min encl. ellipsoid \mathcal{E} of S , with p.s.d. E ;
 set as $\mathcal{E}_{min}, \mathcal{E}_{max}$ the min and max \mathcal{E} axes;
 compute the Cholesky decomposition $L^T L$ of E ;
 transform P and p w.r.t. L ;
until $\mathcal{E}_{max}/\mathcal{E}_{min} < t_r$;
set ρ the largest distance from c to any point in S ;

// MMC
set $\alpha \leftarrow \lfloor \log r \rfloor$; $\beta \leftarrow \lceil \log \rho \rceil$;
 $P_i \leftarrow P \cap B(c, 2^{i/d})$ for $i = \alpha, \alpha + 1, \dots, \beta$;
 $\text{vol}(P_\alpha) \leftarrow 2\pi^{d/2}(2^{\lfloor \log r \rfloor})^d/d\Gamma(d/2)$;
 $i \leftarrow \beta$;
while $i > \alpha$ **do**
 $P_{\text{large}} \leftarrow P_i$; $i \leftarrow i - 1$; $P_{\text{small}} \leftarrow P_i$;
 $\text{count}_{\text{prev}} \leftarrow \text{size}(S)$; remove from S the points not in P_{small} ; $\text{count} \leftarrow \text{size}(S)$;
 Set p to be an arbitrary point from S ;
 for $j = 1$ to $N - \text{count}_{\text{prev}}$ **do**
 $p \leftarrow \text{Walk}(p, P_{\text{large}}, W)$;
 if $p \in B(c, 2^{i/d})$ **then**
 $\text{count} \leftarrow \text{count} + 1$;
 add p in S ;
 $\text{vol} \leftarrow \text{vol} \cdot (N/\text{count})$;
return $\text{vol}/\det(L^T)$;

The following Lemma states the runtime of Alg. 1, which is in fact a variant of the algorithm analysed in [22] (see also Prop. 1). Although there is no theoretical bound on the approximation error of Alg. 1, our experimental analysis in Sect. 4 shows that in practice the achieved error is always better than the one proved in Prop. 1.

Lemma 2. *Given H-polytope P , Alg. 1 performs k phases of rounding in $O^*(d^3mk)$, and approximates $\text{vol}(P)$ in $O(md^3 \log d \log(\rho/r))$ arithmetic operations, assuming $\epsilon > 0$ is fixed, where r and ρ denote the radii of the largest inscribed ball and of the co-centric ball covering P .*

Proof. Our approach generates $d \log(\rho/r)$ balls and uses Hit-and-run. Assuming P contains the unit ball, an upper bound on ρ/r is diameter δ . In each ball intersected with P , we generate $\leq N = 400\epsilon^{-2}d \log d$ random points. Each point is computed after $W = O(d)$ steps of CDHR.

P	d	m	$\text{vol}(P)$	N	μ	[min, max]	std-dev	$\frac{\text{vol}(P)-\mu}{\text{vol}(P)}$	VolEsti (sec)	Exact (sec)
cube-10	10	20	1.024E+003	9210	1.027E+003	[0.950E+003,1.107E+003]	3.16E+001	0.0030	0.42	0.01
cube-15	15	30	3.277E+004	16248	3.24E+004	[3.037E+004,3.436E+004]	9.41E+002	0.0088	1.44	0.40
cube-20	20	40	1.048E+006	23965	1.046E+006	[0.974E+006,1.116E+006]	3.15E+004	0.0028	4.62	swap
cube-50	50	100	1.126E+015	78240	1.125E+015	[1.003E+015,1.253E+015]	4.39E+013	0.0007	117.51	swap
cube-100	100	200	1.268E+030	184206	1.278E+030	[1.165E+030,1.402E+030]	4.82E+028	0.0081	1285.08	swap
Δ -10	10	11	2.756E-007	9210	2.76E-007	[2.50E-007,3.08E-007]	1.08E-008	0.0021	0.56	0.01
Δ -50	60	61	1.202E-082	98264	1.21E-082	[1.07E-082,1.38E-082]	6.44E-084	0.0068	183.12	0.01
Δ -100	100	101	1.072E-158	184206	1.07E-158	[9.95E-159,1.21E-158]	4.24E-160	0.0032	907.52	0.02
Δ -20-20	40	42	1.689E-037	59022	1.70E-037	[1.54E-037,1.87E-037]	7.33E-039	0.0088	53.13	0.01
Δ -40-40	80	82	1.502E-096	140224	1.50E-096	[1.32E-096,1.70E-096]	7.70E-098	0.0015	452.05	0.01
Δ -50-50	100	102	1.081E-129	184206	1.10E-129	[1.01E-129,1.19E-129]	4.65E-131	0.0154	919.01	0.02
cross-10	10	1024	2.822E-004	9210	2.821E-004	[2.693E+004,2.944E+004]	5.15E-006	0.0003	1.58	388.50
cross-11	11	2048	5.131E-005	10550	5.126E-005	[4.888E-005,5.437E-005]	1.15E-006	0.0010	5.19	6141.40
cross-12	12	4096	8.551E-006	11927	8.557E-006	[8.130E-006,9.020E-006]	1.69E-007	0.0007	12.21	—
cross-15	15	32768	2.506E-008	16248	2.505E-008	[2.332E-008,2.622E-008]	5.15E-010	0.0004	541.22	—
cross-18	18	262144	4.09E-011	20810	4.027E-011	[3.97E-011,4.08E-011]	5.58E-013	0.0165	5791.06	—
rh-8-25	8	25	7.859E+002	6654	7.826E+002	[7.47E+002,8.15E+002]	1.93E+001	0.0042	0.30	1.14
rh-8-30	8	30	2.473E+002	6654	2.449E+002	[2.28E+002,2.68E+002]	1.06E+001	0.0099	0.27	5.56
rh-10-25	10	25	5.729E+003	9210	5.806E+003	[5.55E+003,6.06E+003]	1.85E+002	0.0134	0.66	6.88
rh-10-30	10	30	2.015E+003	9210	2.042E+003	[1.96E+003,2.21E+003]	7.06E+001	0.0132	0.67	swap
rv-8-10	8	24	1.409E+019	6654	1.418E+019	[1.339E+019,1.497E+019]	5.24E+017	0.0107	0.37	0.01
rv-8-11	8	54	3.047E+018	6654	3.056E+018	[2.562E+018,3.741E+018]	3.98E+017	0.0028	0.76	0.54
rv-8-12	8	94	4.385E+019	6654	4.426E+019	[4.105E+019,4.632E+019]	2.07E+018	0.0093	0.59	261.37
rv-8-20	8	1191	2.691E+021	6654	2.724E+021	[2.517E+021,2.871E+021]	1.05E+020	0.0123	3.69	swap
rv-8-30	8	4482	7.350E+021	6654	7.402E+021	[7.126E+021,7.997E+021]	2.19E+020	0.0072	12.73	swap
rv-10-12	10	35	2.136E+022	9210	2.155E+022	[1.952E+022,2.430E+022]	1.53E+021	0.0093	1.00	0.01
rv-10-13	10	89	1.632E+023	9210	1.618E+023	[1.514E+023,1.714E+023]	6.23E+021	0.0088	1.24	59.50
rv-10-14	10	177	2.931E+023	9210	2.962E+023	[2.729E+023,3.195E+023]	1.71E+022	0.0135	2.08	swap
cc-8-10	8	70	1.568E+005	26616	1.589E+005	[1.52E+005,1.64E+005]	3.50E+003	0.0138	1.95	0.05
cc-8-11	8	88	1.391E+006	26616	1.387E+006	[1.35E+006,1.43E+006]	2.65E+004	0.0034	2.10	0.08
Fm-4	6	7	8.640E+001	4300	8.593E+001	[7.13E+001,1.12E+002]	8.38E+000	0.0055	0.19	0.01
Fm-5	10	25	7.110E+003	9210	7.116E+003	[6.35E+003,8.10E+003]	3.01E+002	0.0009	0.69	0.02
Fm-6	15	59	2.861E+005	16248	2.850E+005	[2.42E+005,3.22E+005]	1.55E+004	0.0038	3.24	swap
ccp-5	10	56	2.312E+000	9210	2.326E+000	[2.16E+000,2.52E+000]	7.43E-002	0.0064	0.49	38.00
ccp-6	15	368	1.346E+000	16248	1.346E+000	[1.26E+000,1.45E+000]	3.81E-002	0.0002	6.14	swap
\mathcal{B}_8	49	64	4.42E-023	76279	4.46E-023	[4.05E-023,7.32E-024]	1.93E+004	0.0092	192.97	1920.00
\mathcal{B}_9	64	81	2.60E-033	106467	2.58E-033	[2.23E-033,3.07E-033]	2.13E-034	0.0069	499.56	8 days
\mathcal{B}_{10}	81	100	8.78E-046	142380	8.92E-046	[7.97E-046,9.96E-046]	4.99E-047	0.0152	1034.74	6160 days
\mathcal{B}_{11}	100	121	???	184206	1.40E-060	[1.06E-060,1.67E-060]	1.10E-061	???	2398.17	—
\mathcal{B}_{12}	121	144	???	232116	7.85E-078	[6.50E-078,9.31E-078]	5.69E-079	???	4946.42	—
\mathcal{B}_{13}	144	169	???	286261	1.33E-097	[1.13E-097,1.62E-097]	1.09E-098	???	9802.73	—
\mathcal{B}_{14}	169	196	???	346781	5.96E-120	[5.30E-120,6.96E-120]	3.82E-121	???	17257.61	—
\mathcal{B}_{15}	196	225	???	413804	5.70E-145	[5.07E-145,6.52E-145]	1.55E-145	???	31812.67	—

Table 1: Overall results; $\epsilon = 1$, “swap” indicates it ran out of memory and started swapping. “???” indicates that the exact volume is unknown; “—” indicates it didn’t terminate after at least 10h. VINCI is used for exact volume computation except Birkhoff polytopes where birkhoff is used instead.

The boundary oracle of CDHR is implemented in Sect. 2. In particular, k CDHR steps require $O(dm + (k - 1)m + kd)$ arithmetic operations. It holds $d = O(m)$ and $k = \Omega(d)$. Thus, the amortized complexity of a CDHR step is $O(m)$. Overall, the algorithm needs $O(\epsilon^{-2}md^3 \log d \log(\rho/r))$ operations.

Each rounding iteration decreases δ and runs in $O(nd^2(\epsilon^{-1} + \ln d + \ln \ln(n)))$, where n stands for the number of sampled points, and ϵ is the approximation of the minimum volume ellipsoid of Eq. (3). We generate $n = O(d)$ points, each in $O(m)$ arithmetic operations. Hence, rounding runs in $O^*(d^3mk)$, where ϵ is fixed. Moreover, k is typically constant since $k = 1$ is enough to handle, e.g., polytopes with $\rho/r = 100$ in dimension up to 20. \square

Let us check this bound with the experimental data for cubes, products of simplices, and Birkhoff polytopes, with $d \leq 100$ and $\epsilon = 1$, where $m = 2d$, $d + 2$ and $d + 1 + 2\sqrt{d}$, respectively, for the 3 classes, and for cubes $\log(\rho/r) \leq \log(\sqrt{d}) = O(\log d)$. Fig. 2 shows that the 3 classes behave similarly. Performing a fit of $ad^b \log^2 d$, runtime follows $10^{-5}d^{3.08} \log^2 d$ which shows a smaller dependence on d than our bounds, at this range of experiments.

4 Experiments

We implement and experimentally test the above algorithms and methods in the software package VolEsti. The code currently consists of around 2.5K lines in C++ and is open-source¹. It relies on the CGAL library [11] for its d -dimensional kernel to represent objects such as points and

¹<http://sourceforge.net/projects/randgeom>

P	d	ϵ	RDHR				CDHR			
			μ	[min, max]	$(\text{vol}(P) - \mu) / \text{vol}(P)$	VolEsti (sec)	μ	[min, max]	$(\text{vol}(P) - \mu) / \text{vol}(P)$	VolEsti (sec)
\mathcal{B}_5	16	1	2.27E-07	[1.66E-07,2.85E-07]	0.0072	22.90	2.25E-07	[1.87E-07,2.80E-07]	0.0003	4.06
\mathcal{B}_6	25	1	8.53E-13	[3.72E-13,1.22E-12]	0.0982	105.96	9.53E-13	[7.30E-13,1.15E-12]	0.0083	17.26
\mathcal{B}_7	36	1	2.75E-20	[1.78E-21,6.71E-20]	0.4259	479.40	4.82E-20	[3.86E-20,6.18E-20]	0.0056	56.64
cube-10	10	1	1022.8	[944.3951,1103.968]	0.0012	2.03	1026.83	[970.3117,1096.469]	0.0027	0.34
cube-10	10	0.4	—	—	—	—	—	—	0.0011	2.02
cube-20	20	1	1.04E+6	[9.38E+5,1.14E+6]	0.0033	25.44	1.04E+6	[9.74E+5,1.12E+6]	0.0028	4.62

Table 2: Experiments with CDHR vs RDHR; $W = 10$.

d	m	$\text{vol}(P)$	N	μ	[min, max]	std-dev	$(\text{vol}(P) - \mu) / \text{vol}(P)$	VolEsti (sec)	mem. MB	VolEsti* (sec)	mem. MB
10	1024	2.82E-04	9210	2.82E-04	[2.67E-04,3.00E-04]	5.74E-06	0.0001	1.58	35	0.51	42
12	4096	8.55E-06	11927	8.54E-06	[8.04E-06,8.89E-06]	1.72E-07	0.0010	12.21	35	1.62	72
14	16384	1.88E-07	14778	1.88E-07	[1.80E-07,1.99E-07]	4.09E-09	0.0006	237.22	36	6.49	230
16	65536	3.13E-09	17744	3.13E-09	[2.97E-09,3.33E-09]	6.44E-11	0.0004	1430.93	37	32.87	992
18	262144	4.09E-11	20810	4.09E-11	[3.99E-11,4.29E-11]	7.19E-13	0.0013	5791.06	38	188.43	4781

Table 3: Experiments with NN for boundary oracle on cross-polytopes; VolEsti* uses `flann`; $\epsilon = 1$.

vectors, for its LP solver [19], for the approximate minimum ellipsoid [18], and for generating random points in balls. We use `Eigen` [20] for linear algebra. The memory consumption is dominated by the list of random points which needs $O(dN)$ space during the entire execution of the algorithm (Sect. 3). Arithmetic uses the `double` data type of C++, except from the LP solver, which uses the *GNU Multiple Precision arithmetic library* to avoid double exponent overflow. We experimented with several pseudo-random number generators in Boost [29] and chose the fastest, namely mersenne twister generator `mt19937`. All timings are on an Intel Core i5-2400 3.1GHz, 6MB L2 cache, 8GB RAM, 64-bit Debian GNU/Linux.

Data. The following polytopes are tested (the first 7 are from the VINCI webpage):

- cube- d : $\{x = (x_1, \dots, x_d) \mid x_i \leq 1, x_i \geq -1, x_i \in \mathbb{R} \text{ for all } i = 1, \dots, d\}$,
- cross- d : cross polytope, the dual of cube, i.e. $\text{conv}\{-e_i, e_i, i = 1, \dots, d\}$,
- rh- d - m : polytopes constructed by randomly choosing m hyperplanes tangent to the sphere,
- rv- d - n : dual to rh- d - m , i.e. polytopes with n vertices randomly distributed on the sphere,
- cc-8- n : the 8-dimensional product of two 4-dimensional cyclic polyhedra with n vertices,
- ccp- n : complete cut polytopes on n vertices,
- Fm- d : one facet of the metric polytope in dimension d ,
- Δ - d : the d -dimensional simplex $\text{conv}\{e_i, \text{ for } i = 0, 1, \dots, d\}$,
- Δ - d - d : product of two simplices, i.e $\{(p, p') \in \mathbb{R}^{2d} \mid p \in \Delta$ - $d, p' \in \Delta$ - $d\}$,
- skinny-cube- d : $\{x = (x_1, \dots, x_d) \mid x_1 \leq 100, x_1 \geq -100, x_i \leq 1, x_i \geq -1, x_i \in \mathbb{R} \text{ for } i = 2, \dots, d\}$, rotated by 30° in the plane defined by the first two coordinate axes,
- \mathcal{B}_n : the n -Birkhoff polytope (defined below).

Each experiment is repeated 100 times with $\epsilon = 1$ unless otherwise stated. The reported timing for each experiment is the mean of 100 timings. We keep track of and report the *min* and the *max* computed values, the mean μ , and the standard deviation. We measure the accuracy of our method by $(\text{vol}(P) - \mu) / \text{vol}(P)$ and $(\text{max} - \text{min}) / \mu$; unless otherwise stated mean error of approximation refers to the first quantity. The reader should not confuse these quantities which refer to the approximation error that computed *in practice* with ϵ which refers to the *objective* approximation error. Comparing the practical and objective approximation error, our method is in practice more accurate than indicated by the theoretical bounds. In particular, in all experiments all computed values are contained in the interval $((1 - \epsilon)\text{vol}(P), (1 + \epsilon)\text{vol}(P))$, while theoretical results in [22] guarantee only 75% of them. Actually, the above interval is larger than $[\text{min}, \text{max}]$. In general our experimental results show that our software can approximate the volume of general polytopes up to dimension 100 in less than 2 hours with mean approximation error at most 2% (cf. Table 1).

Random walks and oracles. First, we compare the implementations of boundary oracles using membership oracles versus using facet intersection. By performing experiments with RDHR our algorithm approximate the volume of a 10-cube in 42.58 sec using the former, whereas it runs in 2.03 sec using the latter.

We compare RDHR to CDHR. The latter take advantage of more efficient boundary oracle implementations as described in Sect. 2. Table 2 shows that our algorithm using CDHR becomes faster and more accurate than using RDHR by means of smaller $[min, max]$ interval. Additionally, since CDHR is faster we can increase the accuracy (decrease ϵ) and obtain even more accurate results than RDHR, including smaller error $(vol(P) - \mu)/vol(P)$.

Finally, we evaluate our implementations of boundary oracles using *duality* and *NN search* (Sect. 2). The motivation comes from the fact that the boundary oracle becomes slow when the number of facets is large, e.g., for cross- d , $m = 2^d$. We consider state-of-the-art NN software: CGAL’s dD Spatial Searching implements kd-trees [36], ANN [30] implements kd- and BBD-trees, LSH implements Locality Sensitive Hashing [2], and FLANN [31] implements randomized kd-trees. We compare them against our oracle running in $O(m)$, on cross-17, \mathcal{B}_{10} and cpp-7. We build two kd-trees per coordinate, i.e. one per direction, each tree storing the dual of the corresponding lower and upper hulls.

Consider point queries. FLANN, is very fast in high dimensions (typically > 100), but lacks theoretical guarantees. It turns out that `KDTreeSingleIndexParams` on cross- d returns exact results for all ϵ and d tested, since the tree stores vertices of a cube. Compared to the $O(m)$ oracle, for $\epsilon = 0$ it is 10x slower, for $\epsilon = 2$ it is competitive, and for $\epsilon = 5$ it lets us approximate $vol(\text{cross-18})$ with a 40x speed-up, but with extra memory usage (Table 3). On other datasets, FLANN does not always compute the exact NN even for $\epsilon = 0$. ANN, is very fast up to dimension 20 and offers theoretical guarantees. For $\epsilon = 0$, it guarantees the exact NN, but is $> 10^3$ x slower than our $O(m)$ oracle, though it becomes significantly faster for $\epsilon > 1$. In [32], LSH is reported to be 10x slower than FLANN and competitive with ANN, thus we do test it here.

CGAL for point queries is slower than ANN, but can be parametrized to handle hyperplane queries with theoretical guarantees. Given hyperplane H , we set as query point the projection of the origin on H and as distance-function the inner product between points. With the `Sliding_midpoint` rule and $\epsilon = 0$, this is a bit (while ANN is 1000x) slower than our boundary oracle for cross-17. It is important to design methods for which $\epsilon > 0$ accelerates computation so as to use them with approximate boundary oracles.

The above study provides motivation for the design of algorithms that can use approximate boundary queries and hence take advantage of NN software to handle more general polytopes with large number of facets. Of particular relevance is the development of efficient methods and data-structures for approximate hyperplane queries.

Choice of parameters and rounding. We consider two crucial parameters, the length of a random walk, denoted by W , and approximation ϵ , which determines the number N of random points. We set $W = \lfloor 10 + d/10 \rfloor$. Our experiments indicate that, with this choice, either $(vol(P) - \mu)/vol(P)$ or $(min, max)/\mu$ is $< 1\%$ up to $d = 100$ (Table 4). Moreover, for higher W the improvement in accuracy is not significant, which supports the claim that asymptotic bounds are unrealistically high. Fig. 1 correlates runtime (expressed by NW) and accuracy (expressed by $(min, max)/\mu$ which actually measures some “deviation”) to W and ϵ (expressed by N). A positive observation is that accuracy tightly correlates with runtime: e.g., accuracy values close to or beyond 1 lie under the curve $NW = 10^5$, and those rounded to ≤ 0.3 lie roughly above $NW = 3 \cdot 10^5$. It also shows that, increasing W converges faster than increasing N to a value beyond which the improvement in accuracy is not significant.

To experimentally test the effect of *rounding* we construct skinny hypercubes skinny-cube- d . We rotate them to avoid CDHR taking unfair advantage of the degenerate situation where the long edge is parallel to an axis. Table 5 on these and other polytopes shows that rounding reduces approximation error by 2 orders of magnitude. Without rounding, for polytope rv-8-11

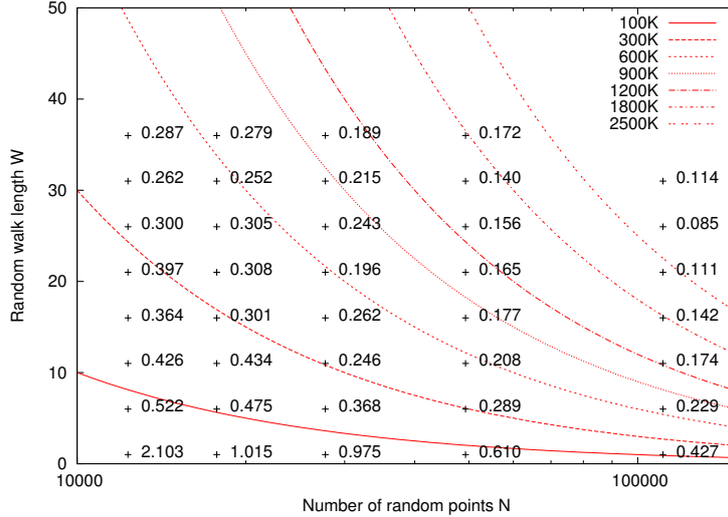


Figure 1: Runtime of `VolEsti` w.r.t. dimension; $\epsilon = 1$, y-axis in logscale; fitting on cube- d results.

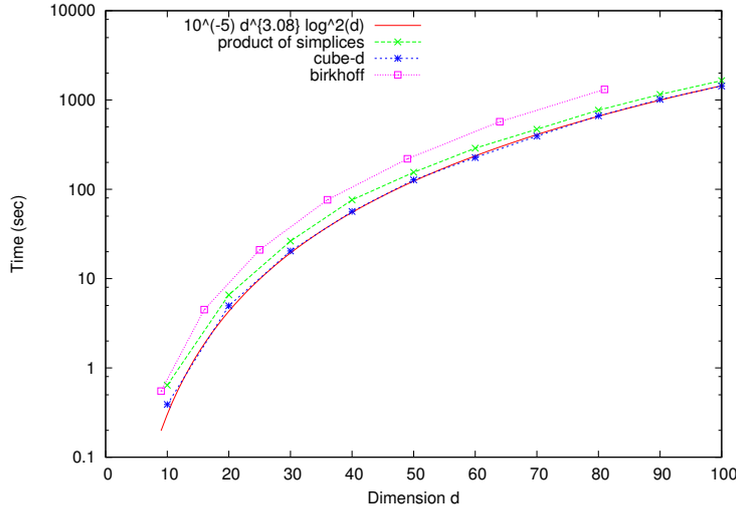


Figure 2: Experiments with \mathcal{B}_5 on the effect of W and ϵ (or N) on accuracy, measured by $(\min, \max)/\mu$ (crosses), and runtime, measured by levels of $N \cdot W = c$, for $c = 10^5, \dots, 2.5 \cdot 10^6$.

one needs to multiply N (thus runtime) by 100 in order to achieve approximation error same as with rounding.

Other software. Exact volume computation concerns software computing the exact value of the volume, up to round-off errors in case it uses floating point arithmetic. We mainly test against VINCI 1.0.5 [8], which implements state-of-the art algorithms, cf. Table 1. For H-polytopes, the method based on Lawrence’s general formula is numerically unstable resulting in wrong results in many examples [9], and thus was excluded. Therefore, we focused on Lasserre’s method. For all polytopes there is a threshold dimension for which VINCI cannot compute the volume: it takes a lot of time (e.g. > 4 hrs for cube-20) and consumes all system memory, thus starts swapping.

LRS is not useful for H-polytopes as stated on its webpage: “If the volume option is applied to an H-representation, the results are not predictable.” `Latte` implements the same decomposition methods as VINCI; it is less prone to round-off error but slower [14]. `Normaliz` applies triangulation: it handles cubes for $d \leq 10$, in < 1 min, but for $d = 15$, it did not terminate after 5 hours. `Qhull` handles V-polytopes but does not terminate for cube-10 nor random polytope

P	d	m	W	μ	$[min,max]$	std-dev	$(\text{vol}(P)-\mu)/\text{vol}(P)$	$(\min, \max)/\mu$
(*) cube-10	10	20	10	1026.953	[925.296,1147.101]	33.91331	0.0029	0.2160
cube-10	10	20	15	1024.157	[928.667,1131.928]	31.34121	0.0002	0.1985
cube-10	10	20	20	1026.910	[932.118,1144.601]	30.97023	0.0028	0.2069
cube-50	50	100	10	1.123E+15	[1.019E+15,1.257E+15]	4.135E+13	0.0022	0.2125
(*) cube-50	50	100	15	1.131E+15	[1.039E+15,1.237E+15]	3.882E+13	0.0044	0.1744
cube-50	50	100	20	1.127E+15	[1.033E+15,1.216E+15]	3.893E+13	0.0007	0.1629
cube-100	100	200	10	1.278E+30	[1.165E+30,1.402E+30]	4.819E+28	0.0081	0.1856
cube-100	100	200	15	1.250E+30	[1.243E+30,1.253E+30]	4.075E+27	0.0140	0.0083
(*) cube-100	100	200	20	1.263E+30	[1.190E+30,1.321E+30]	3.987E+28	0.0038	0.1038
Δ -20-20	40	42	10	1.699E-37	[1.527E-37,1.881E-37]	7.670E-39	0.0056	0.2083
(*) Δ -20-20	40	42	14	1.694E-37	[1.526E-37,1.892E-37]	7.096E-39	0.0025	0.2166
Δ -20-20	40	42	20	1.694E-37	[1.433E-37,1.836E-37]	7.006E-39	0.0024	0.2382
Δ -50-50	100	102	10	1.098E-129	[1.012E-129,1.189E-129]	4.652E-131	0.0154	0.1612
Δ -50-50	100	102	15	1.111E-129	[1.090E-129,1.139E-129]	1.610E-131	0.0281	0.0437
(*) Δ -50-50	100	102	20	1.079E-129	[1.011E-129,1.148E-129]	3.685E-131	0.0015	0.1266
\mathcal{B}_{10}	81	100	10	7.951E-55	[6.291E-55,9.077E-55]	8.533E-56	0.0946	0.3504
\mathcal{B}_{10}	81	100	15	8.124E-55	[7.451E-55,8.774E-55]	5.015E-56	0.0750	0.1629
(*) \mathcal{B}_{10}	81	100	20	7.489E-55	[7.398E-55,7.552E-55]	6.615E-57	0.1472	0.0106

Table 4: Experiments with varying W ; $\epsilon = 1$. (*) indicate minimum W where either $(\text{vol}(P)-\mu)/\text{vol}(P)$ or $(\min, \max)/\mu$ is $< 1\%$.

P	$\text{vol}(P)$	N	μ	$[min,max]$	$\frac{\text{vol}(P)-\mu}{\text{vol}(P)}$	VolEsti(sec)
rv-8-11	3.047E+18	6654	1.595E+18	[6.038E+17,3.467E+18]	0.4766	1.48
rv-8-11	3.047E+18	665421	3.134E+18	[3.134E+18,3.134E+18]	0.0283	157.46
(*) rv-8-11	3.047E+18	6654	3.052E+18	[2.755E+18,3.383E+18]	0.0013	1.34
skinny-cube-10	1.024E+05	9210	5.175E+04	[2.147E+04,1.228E+05]	0.4946	0.69
(*) skinny-cube-10	1.024E+05	9210	1.029E+05	[8.445E+04,1.149E+05]	0.0050	0.71
skinny-cube-20	1.049E+08	23965	4.193E+07	[2.497E+07,7.259E+07]	0.6001	5.59
(*) skinny-cube-20	1.049E+08	23965	1.040E+08	[8.458E+07,1.163E+08]	0.0084	6.70

Table 5: Experiments with rounding; (*): means that we use rounding.

rv-15-60 (Table 6). This should be juxtaposed to the duals, namely our software approximates the volume of cross-10 in 2 sec with $< 1\%$ error and rh-15-60 in 3.44 sec. A general conclusion for exact software is that it cannot handle $d > 15$.

We compare with the most relevant approximation method, namely the `Matlab` implementation of [13] for bodies represented as the intersection of an H-polytope and an ellipsoid. They report that the code is optimized to achieve about 75% success rate for bodies of dimension ≤ 100 and $\epsilon \in [0.1, 0.2]$ (not to be confused with the ϵ of our method). Testing [13] with default options and $\epsilon = 0.1$, our implementation with $\epsilon = 1$ runs faster for $d < 80$, performs roughly 100 times more total Hit-and-run steps and returns significantly more accurate results, e.g. from 4 to 100 times smaller error on cube- d when $d > 70$, and from 5 to 80 times on Birkhoff polytopes (Table 7).

Birkhoff polytopes are well studied in combinatorial geometry and offer an important benchmark. The n -th Birkhoff polytope $\mathcal{B}_n = \{x \in \mathbb{R}^{n \times n} \mid x_{ij} \geq 0, \sum_i x_{ij} = 1, \sum_j x_{ij} = 1, 1 \leq i \leq n\}$, also described as the polytope of the perfect matchings of the complete bipartite graph $K_{n,n}$, the polytope of the $n \times n$ doubly stochastic matrices, and the Newton polytope of the determinant. In [5], they present a complex-analytic method for this volume, implemented in package `birkhoff`, which has managed to compute $\text{vol}(\mathcal{B}_{10})$ in parallel execution, which corresponds to a single processor running at 1 GHz for almost 17 years.

First, $\dim \mathcal{B}_n = n^2 - 2n + 1$: we project \mathcal{B}_n to a subspace of this dimension. Our software, with $\epsilon = 1$, computes the volume of polytopes up to \mathcal{B}_{10} in < 1 hour with mean error of $\leq 2\%$ (Table 1). The computed approximation values improves upon the best known upper bounds

P :	rv-15-				rv-10-				cube-			
	30	40	50	60	100	150	200	250	7	8	9	10
time (sec)	7.7	82.8	473.3	swap	37.3	107.8	282.5	449.0	0.1	2.2	119.5	> 5h

Table 6: Experiments with `qhull`; “swap” indicates it ran out of memory and started swapping; “>5h” indicates it did not terminate after 5 hours.

P	[min, max]	software of [12]				[min, max]	Volesti			
		std-dev	$\frac{\text{vol}(P)-\mu}{\text{vol}(P)}$	# total steps	time(sec)		std-dev	$\frac{\text{vol}(P)-\mu}{\text{vol}(P)}$	# total steps	time(sec)
cube-20	[5.11E+05, 1.55E+06]	1.67E+05	0.0198	7.96E+04	21.48	[9.74E+05, 1.12E+06]	3.15E+04	0.0028	3.61E+06	4.62
cube-30	[6.75E+08, 1.45E+09]	1.72E+08	0.0440	2.22E+05	49.24	[9.91E+08, 1.16E+09]	3.89E+07	0.0039	1.21E+07	17.96
cube-40	[7.90E+11, 1.38E+12]	1.67E+11	0.0731	4.30E+05	88.09	[1.01E+12, 1.23E+12]	4.46E+10	0.0039	2.84E+07	50.72
cube-50	[8.75E+14, 1.45E+15]	1.43E+14	0.0327	7.16E+05	148.06	[1.00E+15, 1.25E+15]	4.39E+13	0.0007	5.49E+07	117.51
cube-60	[8.89E+17, 1.43E+18]	1.64E+17	0.0473	1.15E+06	229.33	[1.06E+18, 1.27E+18]	4.00E+16	0.0051	9.42E+07	222.10
cube-70	[9.01E+20, 1.36E+21]	1.49E+20	0.0707	1.66E+06	427.82	[1.02E+21, 1.32E+21]	5.42E+19	0.0013	1.49E+08	358.93
cube-80	[9.30E+23, 1.36E+24]	1.46E+23	0.1145	2.30E+06	531.46	[1.13E+24, 1.30E+24]	4.42E+22	0.0009	2.21E+08	582.19
cube-90	[1.07E+27, 1.88E+27]	2.20E+26	0.0394	3.30E+06	701.54	[1.09E+27, 1.44E+27]	5.18E+25	0.0019	3.15E+08	875.69
cube-100	[9.53E+29, 1.64E+30]	1.93E+29	0.0357	4.19E+06	884.43	[1.17E+30, 1.40E+30]	4.82E+28	0.0081	4.33E+08	1285.08
\mathcal{B}_8	[2.12E-23, 2.45E-22]	6.25E-23	0.3970	9.31E+05	221.30	[4.05E-23, 7.32E-24]	1.93E+04	0.0092	1.01E+08	192.97
\mathcal{B}_9	[1.54E-33, 2.77E-33]	3.71E-34	0.1830	2.05E+06	420.07	[2.23E-33, 3.07E-33]	2.13E-34	0.0069	2.27E+08	499.56
\mathcal{B}_{10}	[3.39E-46, 1.92E-45]	4.75E-46	0.1207	3.69E+06	691.97	[7.97E-46, 9.96E-46]	4.99E-47	0.0152	4.62E+08	1034.74

Table 7: Comparison of the software [12] vs `VolEsti`; each experiment is run 10 times, total steps refer to the mean of the total number of Hit-and-run steps in each execution.

	n	3	4	5	6	7	8	9	10
$\frac{\text{estimate}}{\text{actual}}$	[10]	1.25408	1.22556	1.19608	1.17258	1.15403	1.13910	1.12684	1.11627
	VolEsti	0.99485	1.09315	1.00029	1.00830	1.00564	0.99440	0.99313	1.01525

Table 8: Comparison between asymptotic and experimental approximation of the volume of \mathcal{B}_n .

on $\text{vol}(\mathcal{B}_n)$, obtained through the asymptotic formula of [10], cf. Table 8. By setting $\epsilon = .5$ we obtain an error of 0.7% for $\text{vol}(\mathcal{B}_{10})$, in 6 hours. The computed approximation of the volume has two correct digits, i.e. its first two digits equal to the ones of the exact volume. More interestingly, using $\epsilon = 1$ we compute, in < 9 hours, an approximation as well as an interval of values for $\text{vol}(\mathcal{B}_{11}), \dots, \text{vol}(\mathcal{B}_{15})$, whose exact values are unknown (Table 1).

5 Further work

NN search seems promising and could accelerate our code, especially if it were performed approximately with hyperplane queries. Producing (almost) uniform point samples is of independent interest in machine learning, including sampling contingency tables and learning the p-value. We plan to exploit such applications of our software. We may also study sampling for special polytopes such as Birkhoff. It is straightforward to parallelize certain aspects of the algorithm, such as random walks assigning each thread to a processor, though other aspects, such as the algorithm’s phases, require more sophisticated parallelization. Our original motivation and ultimate goal is to extend these methods to V-polytopes represented by an optimization oracle.

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