A New Randomized Algorithm to Approximate the Star Discrepancy Based on Threshold Accepting

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Abstract

We present a new algorithm for estimating the star discrepancy of arbitrary point sets. Similar to the algorithm for discrepancy approximation of Winker and Fang [SIAM J. Numer. Anal. 34 (1997), 2028–2042] it is based on the optimization algorithm threshold accepting. Our improvements include, amongst others, a non-uniform sampling strategy which is more suited for higher-dimensional inputs, and rounding steps which transform axis-parallel boxes, on which the discrepancy is to be tested, into critical test boxes. These critical test boxes provably yield higher discrepancy values, and contain the box that exhibits the maximum value of the local discrepancy. We provide comprehensive experiments to test the new algorithm. Our randomized algorithm computes the exact discrepancy frequently in all cases where this can be checked (i.e., where the exact discrepancy of the point set can be computed in feasible time). Most importantly, in higher dimension the new method behaves clearly better than all previously known methods.

1 Introduction

Discrepancy theory analyzes the irregularity of point distributions and has considerable theoretical and practical relevance. There are many different discrepancy notions with a wide range of applications as in optimization, combinatorics, pseudo random number generation, option pricing, computer graphics, and other areas, see, e.g., the monographs [BC87, Cha00, DP10, DT97, FW94, Lem09, Mat09, Nie92, NW10].

In particular for the important task of multivariate or infinite dimensional numerical integration, which arises frequently in fields such as finance, statistics, physics or quantum chemistry, quasi-Monte Carlo algorithms relying on low-discrepancy samples have extensively been studied in

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the last decades. For several classes of integrands the error of quasi-Monte
Carlo approximation can be expressed in terms of the discrepancy of the
set of sample points. This is put into a quantitative form by inequalities of
Koksma-Hlawka- or Zaremba-type, see, e.g., [DP10, Gne11, NW10] and the
literature mentioned therein. The essential point here is that a set of sample
points with small discrepancy results in a small integration error.

Of particular interest are the star discrepancy and the weighted star
discrepancy, which we define below. For theoretical and practical reasons
the weighted star discrepancy attracted more and more attention over the
last few years, see, e.g., [DLP05, HPS08, Joe06, SJ07]. In particular, it is
very promising for finance applications, see [Slo10].

Let \( X = (x^i)_{i=1}^n \) be a finite sequence in the \( d \)-dimensional (half-open)
unit cube \([0,1]^d\). For \( y = (y_1, \ldots, y_d) \in [0,1]^d \) let \( A(y,X) \) be the number
of points of \( X \) lying in the \( d \)-dimensional half-open subinterval \([0,y) := [0,y_1) \times \cdots \times [0,y_d)\), and let \( V_y \) be the \( d \)-dimensional (Lebesgue) volume of
\([0,y)\). We call

\[
d^\ast_\infty(X) := \sup_{y \in [0,1]^d} \left| V_y - \frac{1}{n} A(y,X) \right|
\]

the \( L^\infty \)-star discrepancy, or simply the star discrepancy of \( X \).

For a subset \( u \subseteq \{1, \ldots, d\} \) define \( \Phi_u : [0,1]^d \rightarrow [0,1]^{|u|}, y \mapsto (y_i)_{i \in u} \). For a finite sequence of non-negative weights \((\gamma_u)_{u \subseteq \{1, \ldots, d\}}\) the weighted star
discrepancy of \( X \) is defined by

\[
d^\ast_{\gamma,\infty}(X) := \sup_{0 \neq u \subseteq \{1, \ldots, d\}} \gamma_u d^\ast_\infty(\Phi_u(X)).
\]

Obviously, the star discrepancy is a special instance of the weighted star
discrepancy. Further important discrepancy measures are, e.g., the \( L^p \)-star
discrepancies

\[
d^p_p(X) := \left( \int_{[0,1]^d} \left| V_y - \frac{1}{n} A(y,X) \right|^p \, dy \right)^{1/p}, \quad 1 \leq p < \infty,
\]

and weighted versions thereof. In this article we focus on algorithms to
approximate the star discrepancy, but note that these can be used as bases
for algorithms to approximate the weighted star discrepancy.

In many applications it is of interest to measure the quality of certain sets
by calculating their (weighted or unweighted) star discrepancy, e.g., to test
whether successive pseudo random numbers are statistically independent
[Nie92], or whether given sample sets are suitable for multivariate numerical
integration of certain classes of integrands. As explained in [DGKP08], the
fast calculation or approximation of the (weighted) star discrepancy would
moreover allow efficient randomized semi-constructions of low-discrepancy
samples of moderate size (meaning at most polynomial in the dimension
\( d \)). Actually, there are derandomized algorithms known to construct such
samples deterministically [DGKP08, DGW09, DGW10], but these exhibit high running times. Therefore, efficient semi-constructions would be helpful to avoid the costly derandomization procedures. The critical step in the semi-construction is the efficient calculation (or approximation) of the discrepancy of a randomly chosen set.

The $L^2$-star discrepancy of a given $n$-point set in dimension $d$ can be computed with the help of Warnock’s formula [War72] with $O(dn^2)$ arithmetic operations. Heinrich and Frank provided an asymptotically faster algorithm using $O(n \log n)^{d-1}$ operations for fixed $d$ [FH96, Hei96]. Similarly efficient algorithms are not known for the star discrepancy (and thus also not for the more general weighted star discrepancy). In fact it is known that the problem of calculating the star discrepancy of arbitrary point sets is an $NP$-hard problem [GSW09]. Furthermore, it was shown recently that it is also a $W[1]$-hard problem with respect to the parameter $d$ [GKWW11]. So it is not very surprising that all known algorithms for calculating the star discrepancy or approximating it up to a user-specified error exhibit running times exponential in $d$, see [DEM96, Gne08, Thi01a, Thi01b]. Let us have a closer look at the problem: For a finite sequence $X = \{x_i\}_{i=1}^n$ in $[0, 1)^d$ and for $j \in \{1, \ldots, d\}$ we define

$$\Gamma_j(X) = \{x^j \mid i \in \{1, \ldots, n\}\} \quad \text{and} \quad \bar{\Gamma}_j(X) = \Gamma_j(X) \cup \{1\},$$

and the grids

$$\Gamma(X) = \Gamma_1(X) \times \cdots \times \Gamma_d(X) \quad \text{and} \quad \bar{\Gamma}(X) = \bar{\Gamma}_1(X) \times \cdots \times \bar{\Gamma}_d(X).$$

Then we obtain

$$d^\infty_{\infty}(X) = \max \left\{ \max_{y \in \Gamma(X)} \left( V_y - \frac{1}{n} A(y, X) \right), \max_{y \in \bar{\Gamma}(X)} \left( \frac{1}{n} \bar{A}(y, X) - V_y \right) \right\},$$

(1)

where $\bar{A}(y, X)$ denotes the number of points of $X$ lying in the closed $d$-dimensional subinterval $[0, y]$. (For a proof see [GSW09] or [Nie72, Thm. 2].) Thus, an enumeration algorithm would provide us with the exact value of $d^\infty_{\infty}(X)$. But since the cardinality of the grid $\Gamma(X)$ for almost all $X$ is $n^d$, such an algorithm would be infeasible for large values of $n$ and $d$.

Since no efficient algorithm for the exact calculation or approximation of the star discrepancy up to a user-specified error is likely to exist, other authors tried to deal with this large scale integer programming problem by using optimization heuristics. In [WF97], Winker and Fang used threshold accepting to find lower bounds for the star discrepancy. Threshold accepting [DS90] is a refined randomized local search algorithm based on a similar idea as the simulated annealing algorithm [KGV83]. In [Thi01b], Thiémard gave an integer linear programming formulation for the problem and used techniques as cutting plane generation and branch and bound to tackle it
(cf. also [GSW09]). Quite recently, Shah proposed a genetic algorithm to calculate lower bounds for the star discrepancy [Sha10].

Here in this paper we present a new randomized algorithm to approximate the star discrepancy. As the algorithm of Winker and Fang ours is based on threshold accepting but adds more problem specific knowledge to it. The paper is organized as follows. In Section 2 we describe the algorithm of Winker and Fang. In Section 3 we present a first version of our algorithm. The most important difference to the algorithm of Winker and Fang is a new non-uniform sampling strategy that takes into account the influence of the dimension \( d \) and topological characteristics of the given point set. In Section 4 we introduce the concept of critical test boxes. It are the critical test boxes which lead to the largest discrepancy values, including the maximum value. We present rounding procedures which transform given test boxes into critical test boxes. With the help of these procedures and some other modifications, our algorithm achieves even better results. However, this precision comes at the cost of larger running times (roughly a factor two, see Table 1). In Section 5 we analyze the new sampling strategy and the rounding procedures in more depth. We provide comprehensive numerical tests in Section 6. The results indicate that our new algorithm is superior to all other known methods, especially in higher dimensions. The appendix contains some technical results necessary for our theoretical analyses in Section 5.

2 The Algorithm by Winker and Fang

2.1 Notation

In addition to the notation introduced above, we make use of the following conventions.

For all positive integers \( m \in \mathbb{N} \) we put \([m] := \{1, \ldots, m\}\). If \( r \in \mathbb{R} \), let \([r] := \max\{n \in \mathbb{Z} \mid n \leq r\}\). For the purpose of readability we sometime omit the \([\ ]\) sign, i.e., we whenever we write \( r \) where an integer is required, we implicitly mean \([r]\).

For general \( x, y \in [0, 1]^d \) we write \( x \leq y \) if \( x_j \leq y_j \) for all \( j \in [d] \) and, equivalently, \( x < y \) if \( x_j < y_j \) for all \( j \in [d] \). The characteristic function \( 1_{[0,x)} \) is defined on \([0, 1]^d\) by \( 1_{[0,x)}(y) := 1 \) if \( y < x \) and \( 1_{[0,x)}(y) := 0 \) otherwise. We use corresponding conventions for the closed \( d \)-dimensional box \([0, x]\).

For a given sequence \( X = (x^1)_{i=1}^n \) in the \( d \)-dimensional unit cube \([0, 1)^d\),
we define the following functions. For all $y \in [0, 1]^d$ we set

$$\delta(y) := \delta(y, X) := V_y - A(y, X) = V_y - \frac{1}{n} \sum_{k=1}^{n} 1_{|0, y|}(x^k),$$

$$\bar{\delta}(y) := \bar{\delta}(y, X) := \bar{A}(y, X) - V_y = \frac{1}{n} \sum_{k=1}^{n} 1_{|0, y|}(x^k) - V_y,$$

and $\delta^*(y) := \delta^*(y, X) := \max \{ \delta(y), \bar{\delta}(y) \}$. Then $d^*_\infty(X) = \max_{y \in \Gamma(X)} \delta^*(y)$ as discussed in the introduction.

### 2.2 The algorithm by Winker and Fang

Threshold accepting is an integer optimization heuristic introduced by Dueck and Scheuer in [DS90]. Althöfer and Koschnik [AK91] showed that for suitably chosen parameters, threshold accepting converges to a global optimum if the number $I$ of iterations tends to infinity. Winker and Fang [WF97] applied threshold accepting to compute the star discrepancy of a given $n$-point configuration. In the following, we give a short presentation of their algorithm. A flow diagram of the algorithm can be found in [WF97].

**Initialization:** The heuristic starts with choosing uniformly at random a starting point $x^c \in \bar{\Gamma}(X)$ and calculating $\delta^*(x^c) = \max \{ \delta(x^c), \bar{\delta}(x^c) \}$. Note that throughout the description of the algorithm, $x^c$ denotes the currently used search point.

**Optimization:** A number $I$ of iterations is performed. In the $t$-th iteration, the algorithm chooses a point $x^{nb}$ uniformly at random from a given neighborhood $\mathcal{N}(x^c)$ of $x^c$ and calculates $\delta^*(x^{nb})$. It then computes $\Delta \delta^* := \delta^*(x^{nb}) - \delta^*(x^c)$. If $\Delta \delta^* \geq T$ for a given (non-positive) threshold value $T$, then $x^c$ is being updated, i.e., the algorithm sets $x^c := x^{nb}$. With the help of the non-positive threshold it shall be avoided to get stuck in a bad local maximum $x^c$ of $\delta^*"local"$ with respect to the underlying neighborhood definition. The threshold value $T$ changes during the run of the algorithm and ends up at zero. This should enforce the algorithm to end up at a local maximum of $\delta^*$ which is reasonably close to $d^*_\infty(X)$.

**Neighborhood Structure:** Let us first give the neighborhood definition used in [WF97]. For this purpose, let $x \in \bar{\Gamma}(X)$ be given. Let $\ell < n/2$ be an integer and put $k := 2\ell + 1$. We allow only a certain number of coordinates to change by fixing a value $mc \in [d]$ and choosing $mc$ coordinates $j_1, \ldots, j_{mc} \in [d]$ uniformly at random. For $j \in \{j_1, \ldots, j_{mc}\}$ we consider the set of grid coordinates

$$\mathcal{N}_{k,j}(x) := \left\{ \gamma \in \bar{\Gamma}_j(X) \mid \max\{1, \phi_j^{-1}(x_j) - \ell\} \leq \phi_j^{-1}(\gamma) \leq \min\{|\bar{\Gamma}_j(X)|, \phi_j^{-1}(x_j) + \ell\} \right\},$$

where $\phi_j : |\bar{\Gamma}_j(X)| \rightarrow \bar{\Gamma}_j(X)$ is the ordering of the set $\bar{\Gamma}_j(X)$, i.e., $\phi_j(r) < \phi_j(s)$ for $r < s$. The neighborhood $\mathcal{N}_{k,j_1,\ldots,j_{mc}}(x)$ of $x$ of order $k$ is the
Cartesian product

\[ N_{k,1}^{j_1 \ldots j_{mc}}(x) := \hat{N}_{k,1}(x) \times \ldots \times \hat{N}_{k,d}(x), \]

where \( \hat{N}_{k,j}(x) = \mathcal{N}_{k,j}(x) \) for \( j \in \{j_1, \ldots, j_{mc}\} \) and \( \hat{N}_{k,j}(x) = \{x_j\} \) otherwise. Clearly, \( |N_{k,1}^{j_1 \ldots j_{mc}}(x)| \leq (2^\ell + 1)^{mc} \). We abbreviate \( \mathcal{N}_{mc}^{mc}(x) := \mathcal{N}_{k}^{j_1 \ldots j_{mc}}(x) \) if \( j_1, \ldots, j_{mc} \) are \( mc \) coordinates chosen uniformly at random.

**Threshold values:** Next, we explain how the threshold sequence is chosen in [WF97]. The following procedure is executed prior to the algorithm itself. Let \( I \) be the total number of iterations to be performed by the algorithm and let \( k \leq n \) and \( mc \leq d \) be fixed. For each \( t \in [\sqrt{I}] \), the procedure computes a pair \((y^t, \tilde{y}^t)\), where \( y^t \in \Gamma(X) \) is chosen uniformly at random and \( \tilde{y}^t \in \mathcal{N}_{mc}(y^t) \), again chosen uniformly at random. It then calculates the values \( T(t) := -|\delta^*(y^t) - \delta^*(\tilde{y}^t)| \). When all values \( T(t), t = 1, \ldots, \sqrt{I} \), have been computed, the algorithm sorts them in increasing order. For a given \( \alpha \in (0,1] \), the \( \alpha \sqrt{I} \) values closest to zero are selected as threshold sequence. The number \( J \) of iterations performed for each threshold value is \( J = \alpha^{-1} \sqrt{I} \).

3 A First Improved Algorithm – TA\(_{basic}\)

Our first algorithm, TA\(_{basic}\), builds on the algorithm by Winker and Fang as presented in the previous section. A preliminary but yet different version of TA\(_{basic}\) can be found in [Win07]. It has been used in [DGW10] to provide lower bounds for the comparison of the star discrepancies of different point sequences. In particular in higher dimensions it performed better than any other method tested by the authors.

Recall that the algorithm by Winker and Fang employs a uniform probability distribution on \( \Gamma(X) \) and the neighborhoods \( \mathcal{N}_{mc}(x) \) for all random decisions.

Firstly, this is not appropriate for higher-dimensional inputs: In any dimension \( d \) it is most likely that the discrepancy of a set \( X \) is caused by test boxes with volume at least \( c, c \) some constant in \((0,1)\). Thus in higher dimension \( d \) we expect the upper right corners of test boxes with large local discrepancy to have coordinates at least \( c^{1/d} \). Thus it seems appropriate for higher dimensional sets \( X \) to weight points in the grid \( \Gamma(X) \) with larger coordinates more than points with small coordinates.

Secondly, a uniform probability distribution does not take into account the topological characteristics of the point set \( X \) as, e.g., distances between the points in the grid \( \Gamma(X) \): If there is a grid cell \([x, y]\) in \( \Gamma(X) \) (i.e., \( x, y \in \Gamma(X) \) and \( \phi_j^{-1}(y_j) = \phi_j^{-1}(x_j) + 1 \) for all \( j \in [d] \), where \( \phi_j \) is again the ordering of the set \( \Gamma_j(X) \)) with large volume, we would expect that \( \hat{\delta}(x) \) or \( \delta(y) \) are also rather large.
Thus, on the one hand, it seems better to consider a modified probability measure on $\hat{\Gamma}(X)$ which accounts for the influence of the dimension and the topological characteristics of $X$. On the other hand, if $n$ and $d$ are large, we clearly cannot afford an elaborate precomputation of the modified probability weights.

To cope with this, the non-uniform sampling strategy employed by TA$_\text{basic}$ consists of two steps:

- A continuous sampling step, where we select a point in the whole $d$-dimensional unit cube (or in a “continuous” neighborhood of $x^c$) with respect to a non-uniform (continuous) probability measure $\pi^d$, which is more concentrated in points with larger coordinates.

- A rounding step, where we round the selected point to the grid $\hat{\Gamma}(X)$.

In this way we address both the influence of the dimension and the topological characteristics of the point set $X$. This works without performing any precomputation of probability weights on $\hat{\Gamma}(X)$ – the random generator used, the change of measure on $[0,1]^d$ from the $d$-dimensional Lebesgue measure to $\pi^d$, and our rounding procedure do this implicitly! Theoretical and experimental justifications for our non-uniform sampling strategy can be found in Section 5 and Section 6.

3.1 Sampling of Neighbors

In the following, we present how we modify the probability distribution over the neighborhood sets. Our non-uniform sampling strategy consists of the following two steps.

Continuous Sampling

Consider a point $x \in \hat{\Gamma}(X)$. For fixed $mc \in [d]$ let $j_1, \ldots, j_{mc} \in [d]$ be pairwise different coordinates. For $j \in \{j_1, \ldots, j_{mc}\}$ let $\varphi_j : |\hat{\Gamma}_j(X) \cup \{0\}| \to \hat{\Gamma}_j(X) \cup \{0\}$ be the ordering of the set $\hat{\Gamma}_j(X) \cup \{0\}$ (in particular $\varphi_j(1) = 0$). Let us now consider the real interval $C_{k,j}(x) := [\xi(x_j), \eta(x_j)]$ with

$$\xi(x_j) := \varphi\left(\max\{1, \varphi^{-1}(x_j) - \ell\}\right) \quad \text{and} \quad \eta(x_j) := \varphi\left(\min\{|\hat{\Gamma}_j(X) \cup \{0\}|, \varphi^{-1}(x_j) + \ell\}\right).$$

Our new neighborhood $C_k^{j_1, \ldots, j_{mc}}(x)$ of $x$ of order $k$ is the Cartesian product

$$C_k^{j_1, \ldots, j_{mc}}(x) := \hat{C}_{k,1}(x) \times \ldots \times \hat{C}_{k,d}(x),$$

where $\hat{C}_{k,j}(x) = C_{k,j}(x)$ for $j \in \{j_1, \ldots, j_{mc}\}$ and $\hat{C}_{k,j}(x) = \{x_j\}$ otherwise. We abbreviate $C_k^{mc}(x) := C_k^{j_1, \ldots, j_{mc}}(x)$ if $j_1, \ldots, j_{mc}$ are $mc$ coordinates chosen uniformly at random.

Instead of endowing $C_k^{j_1, \ldots, j_{mc}}(x)$ with the Lebesgue measure on the non-trivial components, we choose a different probability distribution which we
describe in the following. First, let us consider the polynomial product measure
\[ \pi^d(dx) = \otimes_{j=1}^d f(x_j) \lambda(dx_j) \]
with density function \( f : [0, 1] \to \mathbb{R}, r \mapsto dr^{d-1} \)
on \([0, 1]^d\); here \( \lambda = \lambda^1 \) should denote the one-dimensional Lebesgue measure. Notice that in dimension \( d = 1 \) we have \( \pi^1 = \lambda \). Picking a random point \( y \in [0, 1]^d \) with respect to the new probability measure \( \pi^d \) can easily be done in practice by sampling a point \( z \in [0, 1]^d \) with respect to \( \lambda^d \) and then putting \( y := (z_1^{1/d}, \ldots, z_d^{1/d}) \).

We endow \( C_{k,\ldots,j_{mc}}^i(x) \) with the probability distribution induced by the polynomial product measure on the \( mc \) non-trivial components \( C_{k,j_1}(x), \ldots, C_{k,j_{mc}}(x) \). To be more explicit, we map each \( C_{k,j}(x), j \in \{j_1, \ldots, j_{mc}\} \), to the unit interval \([0, 1]\) by
\[ \Psi_j : C_{k,j}(x) \to [0, 1], r \mapsto \frac{r^d - (\xi(x_j))^d}{(\eta(x_j))^d - (\xi(x_j))^d}. \]
Recall that \( \xi(x_j) := \min C_{k,j}(x) \) and \( \eta(x_j) := \max C_{k,j}(x) \). The inverse mapping \( \Psi_j^{-1} \) is then given by
\[ \Psi_j^{-1} : [0, 1] \to C_{k,j}, s \mapsto \left( (\eta(x_j))^d - (\xi(x_j))^d \right)^{1/d} s + (\xi(x_j))^d \].

If we want to sample a random point \( y \in C_{k,\ldots,j_{mc}}^i(x) \), we randomly choose scalars \( s_1, \ldots, s_{mc} \) in \([0, 1]\) with respect to \( \lambda \) and put \( y_j := \Psi_j^{-1}(s_i) \) for \( i = 1, \ldots, mc \). For indices \( j \notin \{j_1, \ldots, j_{mc}\} \) we set \( y_j := x_j \).

**Rounding Procedure:** We round the point \( y \) once up and once down to the nearest points \( y^+ \) and \( y^- \) in \( \Gamma(X) \). More precisely, for all \( j \in [d] \), let \( y^+_j := \min \{x_j^i \in \Gamma_j(X) \mid y_j \leq x_j^i \} \). If \( y_j \geq \min \Gamma_j(X) \) we set \( y^-_j := \max \{x_j^i \in \Gamma_j(X) \mid y_j \geq x_j^i \} \) and in case \( y_j < \min \Gamma_j(X) \), we set \( y^-_j := \max \Gamma_j(X) \).

Obviously, \( A(y^+, X) = A(y, X) \) and thus \( \delta(y^+) = V_{y^+} - A(y^+, X) \geq V_y - A(y, X) = \delta(y) \). Similarly, if \( y_j \geq \min \Gamma_j(X) \) for all \( j \in [d] \), we have \( \bar{A}(y^-, X) = A(y, X) \). Hence, \( \delta(y^-) = \bar{A}(y^-, X) - V_{y^-} \geq \bar{A}(y, X) - V_y = \delta(y) \). If \( y_j < \min \Gamma_j(X) \) for at least one \( j \in [d] \) we have \( \delta(y) \leq 0 \) since \( \bar{A}(y, X) = 0 \). But we also have \( A(y, X) = 0 \) and thus \( \delta^+(y) = \delta(y) \leq \delta(y^+) \). Putting everything together, we have shown that \( \max \{\delta(y^+), \bar{\delta}(y^-)\} \geq \delta^+(y) \).

Since it is only of insignificant additional computational cost to also compute \( \delta(y^-) \) where \( y^+ := y_j^+ = \min \Gamma_j(X) \) and \( y^- := \min \Gamma_j(X) \) for \( j \) with \( y_j < \min \Gamma_j(X) \), we also do that in case at least one such \( j \) with \( y_j < \min \Gamma_j(X) \) exists.

For sampling a neighbor \( x^{nc} \) of \( x^c \) the algorithm thus does the following. First, it samples \( mc \) coordinates \( j_1, \ldots, j_{mc} \in [d] \) uniformly at random. Then it samples a point \( y \in C_{k,\ldots,j_{mc}}^i(x^c) \) as described above, computes the
rounded grid points $y^+, y^-$, and $y^{--}$ and computes the discrepancy $\delta^*_\Gamma(y) := \max\{\delta(y^+), \bar{\delta}(y^-), \bar{\delta}(y^{--})\}$ of the rounded grid points. The subscript $\Gamma$ shall indicate that we consider the rounded grid points. As in the algorithm by Winker and Fang, $\text{TA}_{\text{basic}}$ updates $x^c$ if and only if $\Delta \delta^* = \delta^*_\Gamma(y) - \delta^*(x^c) \geq T$, where $T$ denotes the current threshold. In this case we always update $x^c$ with the best rounded test point, i.e., we update $x^c := y^+$ if $\delta^*_\Gamma(y) = \delta(y^+)$, $x^c := y^-$ if $\delta^*_\Gamma(y) = \bar{\delta}(y^-)$, and $x^c := y^{--}$ otherwise.

### 3.2 Sampling of the Starting Point

Similar to the probability distribution on the neighborhood sets, we sample the starting point $x^c$ as follows. First, we sample a point $x$ from $[0, 1]^d$ according to $\pi^d$. We then round $x$ up and down to $x^+, x^-$, respectively, and again we set $x^c := x^+$ if $\delta^*_{\Gamma}(x) = \delta(x^+)$, $x^c := x^-$ if $\delta^*_{\Gamma}(x) = \bar{\delta}(x^-)$, and we set $x^c := x^{--}$ otherwise.

### 3.3 Computation of Threshold Sequence

The modified neighborhood sampling is also used for computing the sequence of threshold values. If we want the algorithm to perform $I$ iterations, we compute the threshold sequence as follows. For each $t \in \left[\sqrt{I}\right]$ we sample a pair $(y^t, \tilde{y}^t)$, where $y^t \in \bar{\Gamma}(X)$ is sampled as is the starting point and $\tilde{y}^t \in \bar{\Gamma}(X)$ is a neighbor of $y^t$, sampled according to the procedure described in Section 3.1. The thresholds $-|\delta^*(y^t) - \delta^*(\tilde{y}^t)|$ are sorted in increasing order and each threshold will be used for $\sqrt{I}$ iterations of $\text{TA}_{\text{basic}}$. Note that by this choice, we are implicitly setting $\alpha := 1$ in the notion of the algorithm by Winker and Fang.

### 3.4 Choices of $k$ and $mc$

Whereas Winker and Fang have little dependence of the parameter $k = 2\ell + 1$ on the size of the point set (see Section 6), we choose $\ell = \left\lfloor \frac{n}{8} \right\rfloor$, if $n \geq 100$ and $\ell = \left\lfloor \frac{n}{4} \right\rfloor$ otherwise. These settings showed reasonable performance in our experiments. For the number of coordinates that we allow to change, the $mc$ value, we simply use $mc = 2$ throughout since a small value of $mc$ got the best results in [Win07] even for medium- and high-dimensional settings.

### 4 Further Improvements – Algorithm $\text{TA}_{\text{improved}}$

In the following, we present further modifications which we applied to the basic algorithm $\text{TA}_{\text{basic}}$. We call the new, enhanced algorithm $\text{TA}_{\text{improved}}$.

The main improvements, which we describe in more detail below, are the following. (i) A further reduction of the search space by introducing new rounding procedures (“snapping”), (ii) shrinking neighborhoods and
growing number of search directions, and (iii) separate optimization of $\delta$ and $\tilde{\delta}$.

4.1 Further Reduction of the Search Space

We mentioned that for calculating the star discrepancy it is sufficient to test just the points $y \in \bar{\Gamma}(X)$ and to calculate $\delta^*(y)$, cf. equation (1). Therefore $\bar{\Gamma}(X)$ has been the search space we have considered so far. But it is possible to reduce the cardinality of the search space even further.

We obtain the reduction of the search space via a rounding procedure which we call snapping. As this is an important element in the modified algorithm, we now discuss the underlying concept of critical points (or test boxes). For $y \in [0,1]^d$ we define

$$S_j(y) := \prod_{i=1}^{j-1} [0,y_i] \times \{y_j\} \times \prod_{k=j+1}^d [0,y_k], \quad j = 1,\ldots,d.$$ 

We say that $S_j(y)$ is a $\delta(X)$-critical surface if $S_j(y) \cap \{x_1,\ldots,x_n\} \neq \emptyset$ or $y_j = 1$. We call $y$ a $\delta(X)$-critical point if for all $j \in [d]$ the surfaces $S_j(y)$ are critical. Let $C$ denote the set of $\delta(X)$-critical points in $[0,1]^d$.

Let $\tilde{S}_j(y)$ be the closure of $S_j(y)$, i.e.,

$$\tilde{S}_j(y) := \prod_{i=1}^{j-1} [0,y_i] \times \{y_j\} \times \prod_{k=j+1}^d [0,y_k], \quad j = 1,\ldots,d.$$ 

We say $\tilde{S}_j(y)$ is a $\tilde{\delta}(X)$-critical surface if $\tilde{S}_j(y) \cap \{x_1,\ldots,x_n\} \neq \emptyset$. If for all $j \in [d]$ the surfaces $\tilde{S}_j(y)$ are $\tilde{\delta}(X)$-critical, then we call $y$ a $\tilde{\delta}(X)$-critical point. Let $\tilde{C}$ denote the set of $\tilde{\delta}(X)$-critical points in $[0,1]^d$.

We call $y$ a $\delta^*(X)$-critical point if $y \in C^* := C \cup \tilde{C}$.

For $j \in [d]$ let $\nu_j := [\bar{\Gamma}_j(X)]$, and let again $\phi_j : [\nu_j] \to \bar{\Gamma}_j(X)$ denote the ordering of $\bar{\Gamma}_j(X)$. Let $\Phi : [\nu_1] \times \cdots \times [\nu_d] \to \bar{\Gamma}(X)$ be the mapping with components $\phi_j$, $j = 1,\ldots,d$. We say that a multi-index $(i_1,\ldots,i_d) \in [n+1]^d$ is a $\delta(X)$-critical multi-index if $\Phi(i_1,\ldots,i_d)$ is a $\delta(X)$-critical point. We use similar definitions in cases where we deal with $\tilde{\delta}(X)$ or $\delta^*(X)$.

Lemma 4.1. Let $X = \{x_1,\ldots,x_n\}$ be a $n$-point configuration in $[0,1]^d$. Let $C = \mathcal{C}(X)$, $\tilde{C} = \mathcal{C}(X)$, and $C^* = \mathcal{C}^*(X)$ be as defined above. Then $C$, $\tilde{C}$ and $C^*$ are non-empty subsets of $\bar{\Gamma}(X)$. Furthermore,

$$\sup_{y \in [0,1]^d} \delta(y) = \max_{y \in C} \delta(y), \quad \sup_{y \in [0,1]^d} \tilde{\delta}(y) = \max_{y \in \tilde{C}} \tilde{\delta}(y) \quad \text{and} \quad \sup_{y \in [0,1]^d} \delta^*(y) = \max_{y \in C^*} \delta^*(y).$$ 

Proof. The set $C$ is not empty, since it contains the point $(1,\ldots,1)$. Let $y \in C$. By definition, we find for all $j \in [d]$ an index $\sigma(j) \in [n]$ with $y_j = x_j^{\sigma(j)}$ or we have $y_j = 1$. Therefore $y \in \bar{\Gamma}(X)$. Let $z \in [0,1]^d \setminus C$. 

10
Since $\delta(z) = 0$ if $z_j = 0$ for any index $j$, we may assume $z_j > 0$ for all $j$. As $z \notin C$ there exists a $j \in [d]$ where $S_j(z)$ is not $\delta(X)$-critical. In particular, we have $z_j < 1$. Let now $\tau = \hat{\Gamma}_j(X)$ be the smallest value with $z_j < \tau$. Then the point $\hat{z} := (z_1, \ldots, z_{j-1}, \tau, z_{j+1}, \ldots, z_d)$ fulfills $V_{\hat{z}} > V_z$. Furthermore, the sets $[0, \hat{z}) \setminus [0, z)$ and $X$ are disjoint. So $[0, \hat{z})$ and $[0, z)$ contain the same points of $X$. In particular we have $A(\hat{z}, X) = A(z, X)$ and thus, $\delta(\hat{z}) > \delta(z)$. This argument verifies $\sup_{y \in [0,1]^d} \delta(y) = \max_{y \in C} \delta(y)$. The remaining statements of Lemma 4.1 can be proven with similar simple arguments.

We now describe how to use this concept in our algorithm. Let us first describe how we sample a neighbor $x_{\text{nb}}$ of a given point $x^c$. The procedure starts exactly as described in Section 3.1. That is, we first sample $mc$ coordinates $j_1, \ldots, j_{mc} \in [d]$ uniformly at random. Next, we sample $y \in C_{j_1, \ldots, j_{mc}}^{}(x^c)$ according to the probability distribution induced by the polynomial product measure $\pi^d$ on the non-trivial components of $C_{j_1, \ldots, j_{mc}}^{}(x^c)$, cf. Section 3.1. Again we round $y$ up and down to the nearest grid points $y^+, y^-$ and $y^{-\cdot}$, respectively. We then apply the following snapping procedures.

**Snapping down.** We aim at finding a $\tilde{\delta}(X)$-critical point $y_{\text{sn}}^- \leq y^-$ such that the closed box $[0, y_{\text{sn}}^-]^d_{j=1}$ contains exactly the same points of $X$ as the box $[0, y_j^-]^d_{j=1}$. We achieve this by simply setting for all $j \in [d]$

$$y_{\text{sn}}^- := \max\{x_j^i | i \in [n], x^i \in [0, y^-]\}.$$

From the algorithmic perspective, we initialize $y_{\text{sn}}^- := (0, \ldots, 0)$ and check for each index $i \in [n]$ whether $x^i \in [0, y^-]$. If so, we check for all $j \in [d]$ whether $x_j^i \leq y_{\text{sn}}^-$ and update $y_{\text{sn}}^- := x_j^i$ otherwise.

**Snapping up.** Whereas snapping down was an easy task to do, the same is not true for snapping up, i.e., rounding a point to a $\delta(X)$-critical one. More precisely, given a point $y^+$, there are multiple $\delta(X)$-critical points $y_{\text{sn}}^+ \geq y^+$ such that the open box created by $y_{\text{sn}}^+$ contains only those points which are also contained in $[0, y^+)$.

Given that we want to perform only one snapping up procedure per iteration, we use the following random version of snapping upwards. In the beginning, we initialize $y_{\text{sn}}^+ := (1, \ldots, 1)$. Furthermore, we pick a permutation $\sigma$ of $[d]$ uniformly at random from the set $S_d$ of all permutations of set $[d]$. For each point $x \in \{x^i | i \in [n]\}$ we now do the following. If $x \in [0, y^+)$ or $x_j \geq y_{\text{sn}}^+$ for at least one $j \in [d]$, we do nothing. Otherwise

---

1The snapping procedure is the same for $y^-$ and $y^{-\cdot}$. Therefore, we describe it for $y^-$ only.

2Being aware that “snapping up” is an oxymoron, we still use this notation as it easens readability in what follows.
we update $y^{+,\text{sn}}_{\sigma(j)} := x_{\sigma(j)}$ for the smallest $j \in [d]$ with $x_{\sigma(j)} \geq y^{+}_{\sigma(j)}$. After this update, $x$ is no longer inside the open box generated by $y^{+,\text{sn}}$.

Note that snapping up is subject to randomness as the $\delta(X)$-critical point obtained by our snapping procedure can be different for different permutations $\sigma \in S_d$.

The complexity of both snapping procedures is of order $O(nd)$. In our experiments, the snapping procedures caused a delay in the (wall clock) running time by a factor of approximately two, if compared to the running time of $\text{TA}_{\text{basic}}$. It is not difficult to verify the following.

**Lemma 4.2.** Let $X$ be a given $n$-point sequence in $[0, 1]^d$. For all $y \in [0, 1]^d$, the point $y^{+,\text{sn}}$, computed as described above, is $\delta(X)$-critical and both $y^{-,\text{sn}}$ and $y^{-,-,\text{sn}}$ are $\tilde{\delta}(X)$-critical.

In the run of the algorithm we now do the following. Given that we start in some grid point $x^c$, we sample $y \in C^\mu_k(x^c)$ and we round $y$ to the closest grid points $y^+, y^-, y^{-,-} \in \Gamma(X)$ as described in Section 3.1. Next we compute the $\delta(X)$-critical point $y^{+,\text{sn}}$, the $\tilde{\delta}(X)$-critical point $y^{-,\text{sn}}$, and, if $y^- \neq y^{-,-}$, we also compute the $\delta(X)$-critical point $y^{-,-,\text{sn}}$. We decide to update $x^c$ if $\Delta \delta^* = \delta^{*,\text{sn}}(y) - \delta^{*,\text{sn}}(x^c) \geq T$, where $T$ is the current threshold, $\delta^{*,\text{sn}}(y) := \max\{\delta(y^{+,\text{sn}}), \tilde{\delta}(y^{-,\text{sn}}), \tilde{\delta}(y^{-,-,\text{sn}})\}$, and $\delta^{*,\text{sn}}(x^c)$ is the values as was computed in the iteration where $x^c$ was updated last. Note that we do not update $x^c$ with any of the critical points $y^{+,\text{sn}}, y^{-,\text{sn}},$ or $y^{-,-,\text{sn}}$, but only replace $x^c$ with the simple rounded grid points $y^+, y^-, or y^{-,-}$, respectively. More precisely, we update $x^c := y^+$ if $\delta^{*,\text{sn}}(y) = \delta(y^{+,\text{sn}})$, $x^c := y^-$ if $\delta^{*,\text{sn}}(y) = \tilde{\delta}(y^{-,\text{sn}})$, and $x^c := y^{-,-},$ otherwise.

### 4.1.1 Further Variants of the Algorithm

We do not update $x^c$ with the critical points, since our experiments showed that the performance of the algorithm can be significantly improved by updating with the (simply) rounded, not necessarily critical points. This seems to allow the algorithm more flexibility and prevents it from getting stuck in a local optimum too early.

We also tested a variant of the algorithm where we only update the best-so-far solution with $\delta^{*,\text{sn}}(y) := \max\{\delta(y^{+,\text{sn}}), \tilde{\delta}(y^{-,\text{sn}}), \tilde{\delta}(y^{-,-,\text{sn}})\}$ but where all other decisions are only based on the value $\delta^*_c(y) := \max\{\delta(x^c), \tilde{\delta}(y^-)\}$. That is, this algorithm does exactly the same as $\text{TA}_{\text{basic}}$ but in addition computes the $\delta(X)$- and $\tilde{\delta}(X)$-critical points and stores the largest values of $\delta^{*,\text{sn}}$. Clearly, the performance (up to random noise) is better than the one of $\text{TA}_{\text{basic}}$ at the cost of a higher running-time. However, it did not perform as well as the one described above where the decision of whether or not to update a point also depends on the $\delta(X)$- and $\tilde{\delta}(X)$-critical $\delta^{*,\text{sn}}$-values.
4.1.2 Computation of the Starting Point and the Threshold Sequence

When computing the starting point \( x^c \) we first sample a random point \( x \) from \([0,1]^d\) according to \( \pi^d \) (see Section 3.1) and compute \( x^+ \) and \( x^- \), and, if applicable, \( x^{--} \). We also compute the \( \delta(X) \)- and \( \bar{\delta}(X) \)-critical points \( x^{+,sn} \), \( x^{--,sn} \), and \( x^{--,sn} \) and set \( \delta^{*,sn}(x) := \max\{\delta(x^{+,sn}), \bar{\delta}(x^{-,sn}), \bar{\delta}(x^{--,sn})\} \).

We put \( x^c := x^+ \) if \( \delta^{*,sn}(x) = \delta(x^{+,sn}) \), \( x^c := x^- \) if \( \delta^{*,sn}(x) = \bar{\delta}(x^{-,sn}) \), and \( x^c := x^{--} \), otherwise.

For computing the threshold sequence, we also use the \( \delta(X) \)- and \( \bar{\delta}(X) \)-critical \( \delta^{*,sn} \)-values. That is, for \( t = 1, \ldots, \sqrt{T} \) we compute \( t \)-th pair \( (y^t, \bar{y}^t) \) by first sampling a random starting point \( y^t \) as described above (i.e., \( y^t \in \{x^+, x^-, x^{--}\} \) for some \( x \) sampled from \([0,1]^d\) according to \( \pi^d \) and \( y^t = x^+ \) if \( \delta^{*,sn}(x) = \delta(x^{+,sn}) \), \( y^t = x^- \) if \( \delta^{*,sn}(x) = \bar{\delta}(x^{-,sn}) \), and \( y^t = x^{--} \) otherwise). We then compute a neighbor \( \bar{y}^t \in C^{mc}_k(y^t) \) and the maximum of the discrepancy of the \( \delta(X) \)- and \( \bar{\delta}(X) \)-critical points \( \delta^{*,sn}(\bar{y}^t) := \max\{\delta(y^{+,sn}), \bar{\delta}(y^{-,sn}), \bar{\delta}(y^{--,sn})\} \). Finally, we sort the threshold values \( T(t) := -|\delta^{*,sn}(y^t) - \delta^{*,sn}(\bar{y}^t)|, t = 1, \ldots, \sqrt{T} \) in increasing order. This will be our threshold sequence.

4.2 Shrinking Neighborhoods and Growing Number of Search Directions

We add the concept of shrinking neighborhoods, i.e., we consider neighborhoods that decrease in size during the run of the algorithm. The intuition here is the following. In the beginning, we want the algorithm to make large jumps. This allows it to explore different regions of the search space. However, towards the end of the algorithm we want it to become more local, allowing it to explore large parts of the local neighborhood. We implement this idea by iteratively shrinking the \( k \)-value. At the same time, we increase the \( mc \)-value, letting the algorithm explore the local neighborhood more thoroughly.

More precisely, we do the following. In the beginning we set \( \ell := (n-1)/2 \) and \( mc := 2 \). That is, the algorithm is only allowed to change few coordinates of the current search point but at the same time it can make large jumps in these directions. Recall that \( k = 2\ell + 1 \). In the \( t \)-th iteration (out of a total number of \( I \) iterations) we then update

\[
\ell := \frac{n-1}{2} \cdot \frac{I-t}{I} + \frac{t}{I} \quad \text{and} \quad mc := 2 + \frac{t}{I}(d-2).
\]

For the computation of the threshold sequence, we equivalently scale \( k \) and \( mc \) by initializing \( \ell := (n-1)/2 \) and \( mc := 2 \) and then setting for the computation of the \( t \)-th pair \( (y^t, \bar{y}^t) \)

\[
\ell := \frac{n-1}{2} \cdot \frac{\sqrt{T}-t}{\sqrt{T}} + \frac{t}{\sqrt{T}} \quad \text{and} \quad mc := 2 + \frac{t}{\sqrt{T}}(d-2).
\]
Recall that we compute a total number of $\sqrt{T}$ threshold values.

4.3 Separate Optimization of $\delta$ and $\bar{\delta}$

Our last modification is based on the intuition that the star discrepancy is either obtained by an open, subproportionally filled box (i.e., there exists a $y \in \Gamma(X)$ such that $d^*_\infty(X) = \delta(y)$), in which case one might assume that there are many points $\tilde{y}$ with large $\delta(\tilde{y})$-values. Alternatively, if the discrepancy is obtained by a closed, overproportionally filled box (i.e., there exists a $y \in \Gamma(X)$ such that $d^*_\infty(X) = \bar{\delta}(y)$), we assume that there are multiple such points $\tilde{y}$ with large $\bar{\delta}(\tilde{y})$-values. This intuition triggered us to test also the following split variant of the algorithm.

In the $\delta$-version of the algorithm, we only consider open test boxes. That is, whenever we want to sample a random starting point [a random neighbor], we proceed exactly as described in Section 4.1 but instead of computing both $y^+$ and $y^-$ (and, potentially $y^{-,-}$) as well as the $\delta(X)$- and $\bar{\delta}(X)$-critical points $y^{+, sn}$, $y^{-, sn}$, and $y^{-,-, sn}$ in the notation of Section 4.1, we only compute $y^+$ [and $y^{+, sn}$], and we initialize $x^c := y^+$ if and only if $\Delta \delta = \delta(y^{+, sn}) - \delta((x^c)^{+, sn}) \geq T$, where $T$ again denotes the current threshold.

The $\bar{\delta}$-version is symmetric. We compute both $y^-$ and $y^{-,-}$ as well as the $\bar{\delta}(X)$-critical points $y^{-, sn}$ and $y^{-,-, sn}$, and we initialize $x^c := y^-$ [we update $x^c := y^+$ if and only if $\Delta \bar{\delta} = \bar{\delta}(y^{-, sn}) - \bar{\delta}((x^c)^{-, sn}) \geq T]$.

Note that only $\delta$-values (or $\bar{\delta}$-values, respectively) are considered for the computation of the threshold sequence as well.

The algorithm now is the following. We perform $I$ iterations of the $\delta$-version of the algorithm and $I$ iterations of the $\bar{\delta}$-version. The algorithm then outputs the maximum value obtained in either one of the two versions.

It should be noted that a large proportion of the computational cost of TA\_improved lies in the snapping procedures. Thus, running $I$ iterations of the $\delta$-version followed by $I$ iterations of the $\bar{\delta}$-version has a comparable running time to running $I$ iterations of an algorithm of the “mixed” form where we snap each point up and down to the $\delta(X)$- and $\bar{\delta}(X)$-critical grid points. Furthermore, as most modern CPUs are multicore and able to run several programs in parallel, the actual wall-clock cost of switching from TA\_basic to the split version of TA\_improved may be smaller still.

Algorithm 1 summarizes TA\_improved. Note that $\delta(n, d, X, I)$ is the equivalent of Algorithm 2 where we replace $\delta$ by $\bar{\delta}$, $x^+$ by $x^-$ etc. The same is true for Subroutine Thresholds($n, d, X, \sqrt{T}, \bar{\delta}$) for computing the threshold sequence for the $\bar{\delta}$-version.
Algorithm 1: The algorithm \textsc{TA-improved} for computing lower bounds of the star discrepancy $d^*_\infty(X)$.

1 **Input:**
2   Problem instance: $n \in \mathbb{N}$, $d \in \mathbb{N}$, sequence $X = (x^i_{i=1}^n \in [0,1]^d$.
3   Number of iterations $I$.
4 **Computation of a lower bound for $d^*_\infty(X)$:**
5   $\delta := \delta(n, d, X, I)$ /* Output of $I$ iterations of the $\delta$-version.*/
6   $\bar{\delta} := \bar{\delta}(n, d, X, I)$ /* Output of $I$ iterations of the $\bar{\delta}$-version.*/
7 **Output:** $\delta^* := \max\{\delta, \bar{\delta}\}$.

Algorithm 2: The $\delta$-version $\delta(n, d, X, I)$.

1 **Initialization:**
2   $TS = (T(i))_{i=1}^\sqrt{I} := \text{Thresholds}(n, d, X, \sqrt{I}, \delta)$ /*Compute the threshold sequence of length $\sqrt{I}$.*/
3   Sample the starting point: pick $x \in [0,1]^d$ with respect to $\pi^d$ and round $x$ up to the nearest grid point $x^+$.
4   Compute the $\delta(X)$-critical point $x^{+,sn}$ and $\delta(x^{+,sn})$.
5   Initialize $x^c := x^+$, \textsc{global} := $\delta(x^{+,sn})$, \textsc{current} := $\delta(x^{+,sn})$, $\ell := \lfloor \frac{n-1}{2} \cdot \frac{1}{\sqrt{I}} \rfloor$ and $mc := 2$.
6 **for** $i = 1, \ldots, \sqrt{I}$ **do**
7       **for** $t = (i-1)\sqrt{I} + 1, \ldots, (i-1)\sqrt{I} + \sqrt{I}$ **do**
8           Update $\ell := \lfloor \frac{n-1}{2} \cdot \frac{t}{\sqrt{I}} + \frac{1}{2} \rfloor$ and $mc := 2 + \lfloor \frac{1}{d} \rfloor (d-2)$.
9           Sample $y \in C^{mc}_k(x^c)$ as described in Section 3.1.
10          Round $y$ up to the nearest grid point $y^+ \in \Gamma(X)$ and compute the $\delta(X)$-critical point $y^{+,sn}$ as well as $\delta(y^{+,sn})$.
11          if $\delta(y^{+,sn}) > \textsc{global}$ **then** update \textsc{global} := $\delta(y^{+,sn})$.
12          if $\Delta \delta := \delta(y^{+,sn}) - \text{\textsc{current}} \geq T$ **then** update $x^c := y^+$ and \text{\textsc{current}} := $\delta(y^{+,sn})$.

5 Theoretical Analysis

From our main innovations, namely the non-uniform sampling strategy and the rounding procedures “snapping up” and “snapping down”, we already analyzed the snapping procedures and proved that they enlarge the quality of our estimates. The analysis of the non-uniform sampling strategy is much more complicated. One reason is that our sampling strategy strongly interacts with the search heuristic threshold accepting. That is why we confine ourselves to the analysis of the pure non-uniform sampling strategy without considering threshold accepting.
Algorithm 3: Subroutine Thresholds\( (n, d, X, \sqrt{T}, \delta) \) for computing the threshold sequence.

<table>
<thead>
<tr>
<th>1</th>
<th>Initialization:</th>
</tr>
</thead>
<tbody>
<tr>
<td>2</td>
<td>Initialize ( \ell := \left\lfloor \frac{n - 1}{2} \right\rfloor ), and ( mc := 2 ).</td>
</tr>
<tr>
<td>3</td>
<td>for ( t = 1, \ldots, \sqrt{T} ) do</td>
</tr>
<tr>
<td>4</td>
<td>Update ( \ell := \left\lfloor \frac{n - 1}{2} \cdot \sqrt{T} - t \sqrt{T} + t \sqrt{T} \right\rfloor ) and ( mc := 2 + \left\lfloor \frac{t}{\sqrt{T}} \right\rfloor (d - 2) ).</td>
</tr>
<tr>
<td>5</td>
<td>Sample a random point: pick ( x \in [0, 1)^d ) with respect to ( \pi^d ) and round ( x ) up to the nearest grid point ( x^+ ). Compute the ( \delta(X) )-critical point ( x^{+, sn} ) and ( \delta(x^{+, sn}) ).</td>
</tr>
<tr>
<td>6</td>
<td>Sample ( y \in C_k^{mc}(x^+) ) as described in Section 3.1.</td>
</tr>
<tr>
<td>7</td>
<td>Round ( y ) up to the nearest grid point ( y^+ \in \bar{\Gamma}(X) ) and compute the ( \delta(X) )-critical point ( y^{+, sn} ) as well as ( \delta(y^{+, sn}) ).</td>
</tr>
<tr>
<td>8</td>
<td>( \tilde{T}(i) := -</td>
</tr>
<tr>
<td>9</td>
<td>Sort thresholds in increasing order to obtain threshold sequence ( (T(i))_{i=1}^n ) with ( T(i) \leq T(i + 1) ) for all ( i \in [n - 1] ).</td>
</tr>
</tbody>
</table>

In Section 5.1 we prove that sampling in the \( d \)-dimensional unit cube with respect to the probability measure \( \pi^d \) instead of \( \lambda^d \) leads to superior discrepancy estimates. (More precisely, we restrict our analysis for technical reasons to the objective function \( \delta \).) In Section 5.2 we verify that for \( d = 1 \) sampling with respect to the probability distribution induced on \( \bar{\Gamma}(X) \) by sampling with respect to \( \pi^d \) in \([0, 1]^d\) and then rounding to the grid \( \bar{\Gamma}(X) \) leads to better discrepancy estimates than the uniform distribution on \( \bar{\Gamma}(X) \). We comment also on the case \( d \geq 2 \). In Section 5.3 we prove that for random point sets \( X \) the probability of \( x \in \bar{\Gamma}(X) \) being a critical point is essentially an increasing function of the position of its coordinates \( x_j \) in the ordered sets \( \Gamma_j(X), j = 1, \ldots, d \). Recall that critical points yield higher values of the local discrepancy function \( \delta^* \) and include the point that leads to its maximum value. Thus the analysis in Section 5.3 serves as another justification of choosing a probability measure on the neighborhoods which weights points with larger coordinates stronger than points with smaller coordinates.

5.1 Analysis of Random Sampling with Respect to \( \lambda^d \) and \( \pi^d \)

Here we want to show that sampling in the \( d \)-dimensional unit cube with respect to the non-uniform probability measure \( \pi^d \) leads to superior results than sampling with respect to the Lebesgue measure \( \lambda^d \).

Before we start with the theoretical analysis, let us give a strong indication that our non-uniform sampling strategy is much more appropriate in higher dimension than a uniform sampling strategy. In [WF97] Winker and Fang chose in each of the dimensions \( d = 4, 5, 6 \) in a random manner 10
lattice points sets, cf. also our Section 6. They calculated the discrepancy of these sets exactly. If \( \eta_d \) denotes the average value of the coordinates of the points \( y \) with \( \delta^*(y) = \sup_{z \in [0,1]^d} \delta^*(z) \), we get \( \eta_4 = 0.799743 \), \( \eta_5 = 0.840825 \), and \( \eta_6 = 0.873523 \). The expected coordinate value \( \mu_d \) of a point \( x \), randomly sampled from \([0,1]^d\) with respect to the measure \( \pi^d \), is \( \mu_d = d/(d+1) \). So we get \( \mu_4 = 0.8 \), \( \mu_5 = 0.83 \), and \( \mu_6 = 0.857143 \). Note that for using \( \lambda^d \) instead of \( \pi^d \) the expected coordinate value is only 0.5 for all dimensions.

5.1.1 Random Sampling in the Unit Cube with Respect to \( \lambda^d \)

We analyze the setting, where we sample in \([0,1]^d\) with respect to \( \lambda^d \) to maximize the objective function \( \delta \). A similar analysis for \( \delta \) is technically more involved than the proof of Proposition 5.1. Furthermore, it leads to a less clear and also worse result. We comment on this at the end of this subsection.

**Proposition 5.1.** Let \( \varepsilon \in (0,1) \), let \( n, d \in \mathbb{N} \), and let \( X = (x^i)_{i=1}^n \) be a sequence in \([0,1]^d\). Let \( x^* = x^*(X) \in [0,1]^d \) satisfy \( \delta(x^*) = \sup_{x \in [0,1]^d} \delta(x) \).

Let us assume that \( V_{x^*} \geq \varepsilon \). Consider a random point \( r \in [0,1]^d \), sampled with respect to the probability measure \( \lambda^d \). If \( P^\lambda_{\varepsilon} = P^\lambda_{\varepsilon}(X) \) denotes the probability of the event \( \{ r \in [0,1]^d \mid \delta(x^*) - \delta(r) \leq \varepsilon \} \), then

\[
P^\lambda_{\varepsilon} \geq \frac{1}{d!} \frac{\varepsilon^d}{V_{x^*}^{d-1}} \geq \frac{\varepsilon^d}{d!}.
\]

This lower bound is sharp in the sense that there exist sequences of point configurations \( \{X^{(k)}\} \) such that \( \lim_{k \to \infty} d! \varepsilon^{-d} P^\lambda_{\varepsilon}(X^{(k)}) \) converges to 1 as \( \varepsilon \) tends to zero.

Let additionally \( \varepsilon \in (0,1) \) and \( R \in \mathbb{N} \). Consider random points \( r^1, \ldots, r^R \in [0,1]^d \), sampled independently with respect to \( \lambda^d \), and put \( \delta^R := \max_{i=1}^R \delta(r^i) \). If

\[
R \geq \| \ln(\varepsilon) \| \ln \left( 1 - \frac{\varepsilon^d}{d!} \right)^{-1},
\]

then \( \delta(x^*) - \delta^R \leq \varepsilon \) with probability at least \( 1 - \varepsilon \).

Notice, that the case \( V_{x^*} \leq \varepsilon \) left out in Proposition 5.1 is less important for us, since our main goal is to find a good lower bound for the star-discrepancy \( d^*_\infty(X) \). Indeed, the approximation of \( d^*_\infty(X) \) up to an admissible error \( \varepsilon \) is a trivial task if \( d^*_\infty(X) \leq \varepsilon \). If \( d^*_\infty(X) > \varepsilon \), then \( V_{x^*} < \varepsilon \) implies \( \delta(x^*) < d^*_\infty(X) \), and the function \( \delta \) plays the significant role.

**Proof.** For \( x \leq x^* \) we get

\[
\delta(x) = V_x - \frac{1}{n} \sum_{i=1}^n 1_{[0,x]}(x^i) \geq \delta(x^*) - (V_{x^*} - V_x).
\]
Therefore the Lebesgue measure of the set
\[ A_{\varepsilon}(x^*) := \{ x \in [0,1]^d \mid x \leq x^*, V_{x^*} - V_x \leq \varepsilon \} \]  
(6)
is a lower bound for \( P_{\varepsilon}^{\lambda} \). Due to Proposition A.2, we have for \( d \geq 2 \)
\[ \lambda^d(A_{\varepsilon}(x^*)) = \frac{1}{d!} \frac{\varepsilon^d}{V_{x^*}^{d-1}} \sum_{k=0}^{\infty} b_k(d) \left( \frac{\varepsilon}{V_{x^*}} \right)^{k}, \]
with positive coefficients \( b_k(d) \). In particular, we have \( b_0(d) = 1 \). Thus,
\[ \lambda^d(A_{\varepsilon}(x^*)) \geq \frac{1}{d!} \frac{\varepsilon^d}{V_{x^*}^{d-1}} \geq \frac{\varepsilon^d}{d!}, \]  
(7)and this estimate is obviously also true for \( d = 1 \). Let us now consider for sufficiently large \( k \in \mathbb{N} \) point configurations \( X^{(k)} = (x^{(k),i})_{i=1}^{n} \), where
\[ x^{(k),1}_1 = \ldots = x^{(k),n}_1 = k/(k+1) > \varepsilon \]  
(8)and \( x^{(k),i}_j < k/(k+1) - \varepsilon \) for all \( i \in [n], j > 1 \). Then obviously \( x^*(X^{(k)}) = (k/(k+1), 1, \ldots, 1) \), and it is easy to see that \( P_{\varepsilon}^{\lambda}(X^{(k)}) = \lambda^d(A_{\varepsilon}(x^*)) \). From Proposition A.2 we get
\[ \lambda^d(A_{\varepsilon}(x^*)) = \left( \frac{k+1}{k} \right)^{d-1} \frac{\varepsilon^d}{d!} \left( 1 + O\left( \frac{k+1}{k} \varepsilon \right) \right). \]
This proves that estimate (4) is sharp. Notice that we assumed (8) only for simplicity. Since \( \delta(x^*(X)) \) is continuous in \( X \), we can find for fixed \( k \) an open set of point configurations doing essentially the same job as \( X^{(k)} \).

Assume now \( \delta(x^*) - \delta^R > \varepsilon \), i.e., \( \delta(x^*) - \delta(r^i) > \varepsilon \) for all \( i \leq R \). The probability of this event is at maximum \( (1 - \varepsilon^d/d!)^R \). This probability is bounded from above by \( \varepsilon \) if \( R \) satisfies (5).

For \( d \geq 1 \) and \( \varepsilon \leq 1/2 \) we have \( \left| \ln(1 - \varepsilon^d/d!) \right|^{-1} \sim d! \varepsilon^{-d} \). In this case we can only assure that \( \delta^R \) is an \( \varepsilon \)-approximation of \( \sup_{x \in [0,1]^d} \delta(x) \) with a certain probability if the number \( R \) of randomly sampled points is super-exponential in \( d \).

Let us end this section with some comments on the setting where we are only interested in maximizing \( \delta \). If for given \( \varepsilon > 0 \), \( X \in [0,1]^n \) the maximum of \( \delta \) is achieved in \( \bar{x} = \bar{x}(X) \in [0,1]^d \), and if we want to know the probability of the event \( \{ r \in [0,1]^d \mid \delta(\bar{x}) - \delta(r) \leq \varepsilon \} \), there seems to be no alternative to estimating \( \lambda^d(U(\bar{x})) \), where
\[ U(\bar{x}) := \{ r \in [0,1]^d \mid \bar{x} \leq r, V_r - V_{\bar{x}} \leq \varepsilon \}. \]
It is easy to see that \( \lambda^d(U(\bar{x}(X))) \) approaches zero if one of the coordinates of \( \bar{x} \) tends to 1 - regardless of \( \varepsilon \) and \( V_{\bar{x}} \). We omit a tedious error analysis to cover the \( \tilde{\delta} \)-setting.
5.1.2 Random Sampling in the Unit Cube with Respect to $\pi^d$

Similarly as in the preceding section, we analyze here the setting where, in order to maximize $\delta$, we sample in $[0,1]^d$ with respect to $\pi^d$.

**Proposition 5.2.** Let $\varepsilon, d, n, X = (x^i)_{i=1}^n$ and $x^*$ as in Proposition 5.1. Again assume $V_{\varepsilon*} \geq \varepsilon$. Consider a random point $r \in [0,1]^d$, sampled with respect to the probability measure $\pi^d$. If $P^\pi_r = P^\pi_r(X)$ denotes the probability of the event $\{ r \in [0,1]^d \mid \delta(x^*) - \delta(r) \leq \varepsilon \}$, then $P^\pi_r \geq \varepsilon^d$. This lower bound is sharp, since there exists a point configuration $X$ such that $P^\pi_r(X) = \varepsilon^d$.

Let additionally $\varepsilon \in (0,1)$ and $R \in \mathbb{N}$. Consider random points $r^1, \ldots, r^R \in [0,1]^d$, sampled independently with respect to $\pi^d$, and put $\delta^R := \max_{i=1}^R \delta(r^i)$. If

$$R \geq |\ln(\varepsilon)| |\ln(1 - \varepsilon^d)|^{-1},$$

then $\delta(x^*) - \delta^R \leq \varepsilon$ with probability at least $1 - \varepsilon$.

**Proof.** Clearly $P^\pi_r \geq \pi^d(A_\varepsilon(x^*))$, where $A_\varepsilon(x^*)$ is defined as in (6). Due to Proposition A.5 we have $\pi^d(A_\varepsilon(x^*)) \geq \varepsilon^d$. Let us now consider the point configuration $X$, where $x^1_1 = \ldots = x^n_1 = \varepsilon$ and $x^i_j < \varepsilon$ for all $i \in [n], j > 1$. Furthermore, at least an $\varepsilon^{-1}$-fraction of the points should be equal to $(\varepsilon, 0, \ldots, 0)$. Then obviously $x^*(X) = (\varepsilon, 1, \ldots, 1)$ and $P^\lambda_r(X) = \pi^d(A_\varepsilon(x^*)) = \varepsilon^d$.

Let us now assume that $\delta(x^*) - \delta^R > \varepsilon$, i.e., $\delta(x^*) - \delta(r_i) > \varepsilon$ for all $i \leq R$. This happens with probability not larger than $(1 - \varepsilon^d)^R$. Therefore we have $(1 - \varepsilon^d)^R \leq \varepsilon$ if $R$ satisfies (9).

If $d \geq 1$ and $\varepsilon \leq 1/2$, then $|\ln(1 - \varepsilon^d)|^{-1} \sim \varepsilon^{-d}$. Here the number of iterations $R$ ensuring with a certain probability that $\delta^R$ is an $\varepsilon$-approximation of $\sup\{|\delta(x)| \mid x \in [0,1]^d\}$ is still exponential in $d$, but at least not super-exponential as in the previous section.

 Altogether we see that a simple sampling algorithm relying on the probabilistic measure $\pi^d$ rather than on $\lambda^d$ is more likely to find larger values of $\delta$.

5.2 Analysis of Rounding to the Coordinate Grid

As described in Sections 3.1 and 3.2, our non-uniform sampling strategy on the grids $\Gamma(X)$ and $\Gamma(X)$ for the objective functions $\delta$ and $\delta$ consists of sampling in $[0,1]^d$ with respect to $\pi^d$ and then rounding the sampled point $y$ up and down to grid points $y^+$ and $y^-$, respectively. This induces discrete probability distributions $w_u = (w_u(z))_{z \in \Gamma(X)}$ and $w_l = (w_l(z))_{z \in \Gamma(X)}$ on $\Gamma(X)$ and $\Gamma(X)$, respectively. If we use additionally the rounding procedures “snapping up” and “snapping down”, as described in Section 4.1, this will lead to modified probabilistic distributions $w_{u\text{sn}} = (w_{u\text{sn}}(z))_{z \in \Gamma(X)}$
and \(w_i^{sn} = (w_i^{sn}(z))_{z \in \Gamma(X)}\) on \(\hat{\Gamma}(X)\) and \(\Gamma(X)\), respectively. In dimension \(d = 1\) the probability distributions \(w_u\) and \(w_u^{sn}\) as well as \(w_i\) and \(w_i^{sn}\) are equal, since every test box is a critical one. Essentially we prove in the next section that in the one-dimensional case sampling with respect to the probability distributions \(w_u = w_u^{sn}\) leads to larger values of \(\delta[\hat{\delta}]\) than sampling with respect to the uniform distribution on \(\hat{\Gamma}(X)\) [\(\Gamma(X)\)].

5.2.1 Analysis of the 1-Dimensional Situation

Recall that in the 1-dimensional case \(\pi = \pi^1\) coincides with \(\lambda = \lambda^1\).

To analyze the 1-dimensional situation, let \(X := (x^i)_{i=1}^n\) be the given point configuration in \([0, 1]\). Without loss of generality we assume that \(0 \leq x^1 < \cdots < x^n < 1\). Since \(\delta^*(1) = 0\) we do not need to consider the whole grid \(\hat{\Gamma}(X)\) but can restrict ourselves to the set \(\Gamma(X) = \{x^1, \ldots, x^n\}\). For the same reason, let us set \(y^+ := x^1\) if \(y > x^n\) (recall that, following the description given in Section 3.1, we set \(y^- := x^n\) for \(y < x^1\) anyhow).

As discussed above, we take points randomly from \(\Gamma(X)\), but instead of using equal probability weights on \(\Gamma(X)\), we use the probability distributions \(w_u = w_u^{sn}\) and \(w_i = w_i^{sn}\) on \(\Gamma(X)\) to maximize our objective functions \(\delta\) and \(\hat{\delta}\), respectively. If we put \(x^0 := x^n - 1\) and \(x^{n+1} := x^1 + 1\), then the corresponding probability weights for \(\delta\) and \(\hat{\delta}\) are given by \(w_i(x^i) := x^i - x^{i-1}\) and \(w_u(x^i) := x^{i+1} - x^i\), respectively.

In the next lemma we will prove the following statements rigorously: If one wants to sample a point \(\tau \in \Gamma(X)\) with \(\delta(\tau)\) as large as possible or if one wants to enlarge the chances to sample the point \(\tau\) where \(\delta\) takes its maximum, its preferable to use the weights \(w_i\) instead of the equal weights \(1/n\) on \(\Gamma(X)\). Similarly, it is preferable to employ the weights \(w_u(x^i), i = 1, \ldots, n\), instead of equal weights if one wants to increase the expectation of \(\delta\) or the chances of sampling the maximum of \(\hat{\delta}\).

**Lemma 5.3.** Let \(d = 1\) and \(\tau, \hat{\tau} \in \Gamma(X)\) with \(\delta(\tau) = \sup_{z \in [0, 1]} \delta(z)\) and \(\hat{\delta}(\hat{\tau}) = \sup_{z \in [0, 1]} \hat{\delta}(z)\). Then we have \(w_i(\tau) \geq 1/n\) and \(w_u(\tau) \geq 1/n\).

Furthermore, let \(\mathbb{E}, \mathbb{E}_l,\) and \(\mathbb{E}_u\) denote the expectations with respect to the uniform weights \(\{1/n\}\), the weights \(\{w_i(x^i)\}\), and the weights \(\{w_u(x^i)\}\) on the probability space \(\Gamma(X)\), respectively. Then \(\mathbb{E}_l(\delta) \geq \mathbb{E}(\delta)\) and \(\mathbb{E}_u(\hat{\delta}) \geq \mathbb{E}(\hat{\delta})\).

**Proof.** Let \(\nu \in [n]\) with \(\tau = x^\nu\). Assume first \(w_i(x^\nu) < 1/n\), i.e., \(x^\nu - x^{\nu - 1} < 1/n\). If \(\nu > 1\), then

\[
\delta(x^{\nu - 1}) = x^{\nu - 1} - \frac{\nu - 2}{n} > x^\nu - \frac{\nu - 1}{n} = \delta(x^\nu).
\]

If \(\nu = 1\), then however

\[
\delta(x^n) = x^n - \frac{n - 1}{n} = x^0 + \frac{1}{n} > x^1 = \delta(x^1).
\]
So both cases result in a contradiction.

We prove now \( \mathbb{E}_l(\delta) \geq \mathbb{E}(\delta) \) by induction over the cardinality \( n \) of \( X \). For \( n = 1 \) we trivially have \( \mathbb{E}_l(\delta) = x^1 = \mathbb{E}(\delta) \).

Therefore, let the statement be true for \( n \) and consider an ordered set \( \Gamma(X) := \{x^1, \ldots, x^{n+1}\} \). Let \( \delta \) achieve its maximum in \( x^\nu \in \Gamma(X) \). We already proved that \( w_l(x^\nu) = x^\nu - x^{\nu-1} \geq 1/(n + 1) \) holds. With the notation

\[
\tilde{x}^i := x^i \text{ if } 1 \leq i < \nu, \quad \tilde{x}^i := x^{i+1} - \frac{1}{n + 1} \text{ if } i \geq \nu,
\]

and

\[
\hat{x}^i := \frac{n + 1}{n} \tilde{x}^i, \quad i = 1, \ldots, n, \quad \text{and} \quad x^0 := \hat{x}^n - 1,
\]

we get

\[
\mathbb{E}_l(\delta) = \sum_{i=1}^{n+1} w_l(x^i)\delta(x^i) = \frac{\delta(x^\nu)}{n + 1} + \left( w_l(x^\nu) - \frac{1}{n + 1} \right) \delta(x^\nu) + \sum_{i=1}^{n+1} w_l(x^i)\delta(x^i) \geq \frac{\delta(x^\nu)}{n + 1} + \left( w_l(x^{\nu+1}) + w_l(x^\nu) - \frac{1}{n + 1} \right) \delta(x^{\nu+1}) + \sum_{i \notin \{\nu, \nu+1\}} w_l(x^i)\delta(x^i) = \frac{\delta(x^\nu)}{n + 1} + \left( \tilde{x}^1 - \tilde{x}^n + \frac{n}{n + 1} \right) \tilde{x}^1 + \sum_{i=2}^{n} (\tilde{x}^i - \tilde{x}^{i-1}) \left( \tilde{x}^i - \frac{i - 1}{n + 1} \right) = \frac{\delta(x^\nu)}{n + 1} + \left( \frac{n}{n + 1} \right)^2 \sum_{i=1}^{n} (\tilde{x}^i - \tilde{x}^{i-1}) \left( \tilde{x}^i - \frac{i - 1}{n} \right).
\]

On the other hand we have

\[
\mathbb{E}(\delta) = \sum_{i=1}^{n+1} \frac{1}{n + 1} \delta(x^i) = \frac{\delta(x^\nu)}{n + 1} + \sum_{i=1}^{n} \frac{1}{n + 1} \left( \tilde{x}^i - \frac{i - 1}{n + 1} \right) = \frac{\delta(x^\nu)}{n + 1} + \left( \frac{n}{n + 1} \right)^2 \sum_{i=1}^{n} \frac{1}{n} \left( \tilde{x}^i - \frac{i - 1}{n} \right).
\]

These calculations and our induction hypothesis, applied to \( \{\tilde{x}^1, \ldots, \tilde{x}^n\} \), lead to \( \mathbb{E}_l(\delta) \geq \mathbb{E}(\delta) \). For \( \mu \in [n] \) with \( \bar{\tau} = x^\mu \) the inequalities \( w_{\mu}(x^\mu) \geq 1/n \) and \( \mathbb{E}_{\mu}(\delta) \geq \mathbb{E}(\delta) \) can be proved in a similar manner.

**5.2.2 Analysis of Higher Dimensional Situations** \( d \geq 2 \)

If we turn to the situation in dimension \( d \geq 2 \), we first face a technical difference: For many sequences \( X \) the supremum \( \sup_{x \in [0,1]^d} \delta(x) \) will be achieved by some \( x \in \Gamma(X) \) with \( x_j = 1 \notin \{x^1_j, \ldots, x^n_j\} \) for at least one
Figure 1: $\delta$ takes its maximum in $x^3$, but $w_l(x^3) < 1/36$
5.3 in $d \geq 2$. More precisely: If $\tau \in \Gamma(X)$ and $\bar{\tau} \in \Gamma(X)$ are points with
\[ \delta(\tau) = \sup_{y \in [0,1]^d} \delta(y) \text{ and } \bar{\delta}(\bar{\tau}) = \sup_{y \in [0,1]^d} \bar{\delta}(y), \]
then in general we do not have
\[ w_l(\tau) \geq \prod_{j=1}^{d} \frac{1}{\nu_j} \text{ and } w_u(\bar{\tau}) \geq \prod_{j=1}^{d} \frac{1}{\nu_j - 1}, \]
even $w_l(\tau)$ or $w_u(\bar{\tau}) \geq (n+1)^{-d}$ need not to be satisfied. Notice that for fixed
\[ d \text{ and } n \]
the sets of counterexamples to each of the inequalities in (10) have
strictly positive measure. Indeed, the suprema of $\delta$ and $\bar{\delta}$ are continuous in
$X$, and the weights $w_l$, $w_u$ are continuous in $X$ as long as for each $j \in [d]$ all
coordinates $x^1_j, \ldots, x^n_j$ are distinct. But it is, e.g., easy to rearrange the
counterexamples in Figure 1 and 2 slightly such that these constraints are
satisfied.

But notice also that in both figures the grid point $\gamma$ provides a good
approximation of the actual maximum of $\delta$ and $\bar{\delta}$, and $\gamma$ has a rather large
weight $w_l$ and $w_u$, respectively. Furthermore, the point sets in Figure 1 and 2
are quite artificial and also far away from being well-distributed. For random
or low-discrepancy sets these probability weights can still be very useful –
this is also confirmed by our numerical experiments, see Section 6. Moreover,
the invalidity of (10) does not necessarily mean that the expectations $E(\delta)$
and $E(\bar{\delta})$ are larger than $E_l(\delta)$ and $E_u(\bar{\delta})$, respectively.

Actually, if we use additionally the procedures “snapping up” and “snap-
ing down” and look at the induced probability distribution $w^{sn}_l$ and $w^{sn}_u$,
respectively, then $x^3$ has the maximum weights $w^{sn}_l$ of all critical points in
Figure 1, and $x^2$ has the maximum weight $w^{sm}_n$ of all critical points in Figure 2. Thus adding the snapping procedures can change the situation decisively.

Additional theoretical results in this direction would be interesting.

5.3 The Chances of a Grid Point Being a Critical Point

If $X = (x^1, \ldots, x^n)$ is a sequence in $[0,1]^d$ which has been chosen randomly with respect to the Lebesgue measure and if $x \in \Gamma(X)$, then the larger the components of $x$, the higher is the probability of $x$ being a $\delta(X)$-critical point. The same holds for $\bar{x} \in \Gamma(X)$ and $\delta(X)$, respectively.

**Proposition 5.4.** Consider $[0,1]^{nd}$ as a probability space endowed with the probability measure $\lambda^{nd}$. Let $\iota := (i_1, \ldots, i_d) \in [n+1]^d$. If $k$ indices $i_{\nu(1)}, \ldots, i_{\nu(k)}$ of the $i_j$, $j = 1, \ldots, d$, are at most $n$ and the remaining $d-k$ of them are equal to $n+1$, then for uniformly distributed random variable $X$ in $[0,1]^{nd}$ the multi-index $\iota$ is $\delta(X)$-critical with probability

$$
\left( \frac{(n-k)!}{n!} \right)^{k-1} \frac{k-1}{k} \prod_{j=1}^{k-1} \max\{i_{\nu(j)} - \ell, 0\}.
$$

**Proof.** Let $\Phi = (\phi_1, \ldots, \phi_d)$ be as in Section 4.1. Since the event that for all coordinates $j \in [d]$ we have $|\Gamma_j(X)| = n+1$ holds with probability 1, we restrict ourselves to this situation.

Without loss of generality, we may assume that $i_1, \ldots, i_k \leq n$ and $i_{k+1} = \ldots = i_d = n+1$. Obviously, $S_{k+1}(\Phi(\iota)), \ldots, S_d(\Phi(\iota))$ are $\delta$-critical surfaces, since $\phi_{k+1}(n+1) = \ldots = \phi_d(n+1) = 1$. For $i = 1, \ldots, k$ let $\sigma_i = \sigma_i(X) : [n] \to [n]$ be the permutation with

$$
x_i^{\sigma_i(1)} < x_i^{\sigma_i(2)} < \cdots < x_i^{\sigma_i(n)} < 1.
$$

Clearly, $\Phi(\iota)_j = \phi_j(i_j) = \sigma_j^{\sigma(i_j)}$ for all $j \in [k]$. Since $S_i(x) \cap S_j(x) = \emptyset$ for all $i \neq j$ and all $x \in [0,1]^d$, the surfaces $S_1(\Phi(\iota)), \ldots, S_k(\Phi(\iota))$ can only be $\delta(X)$-critical if $\{\sigma_1(i_1), \ldots, \sigma_k(i_k)\} = k$. More precisely, $\iota$ is a $\delta(X)$-critical multi-index if and only if the condition

$$
\forall j \in [k] \forall l \in [k] \setminus \{j\} : x_l^{\sigma_l(i_j)} < \Phi(\iota)_l = x_l^{\sigma_l(i_l)}
$$

holds. This is equivalent to the $k$ conditions

$$
\sigma_1^{-1}(\sigma_2(i_2)), \sigma_3^{-1}(\sigma_3(i_3)), \ldots, \sigma_1^{-1}(\sigma_k(i_k)) < i_1, \\
\sigma_2^{-1}(\sigma_1(i_1)), \sigma_3^{-1}(\sigma_3(i_3)), \ldots, \sigma_2^{-1}(\sigma_k(i_k)) < i_2, \\
\vdots \quad \vdots \quad \vdots \quad \vdots \\
\sigma_k^{-1}(\sigma_1(i_1)), \sigma_k^{-1}(\sigma_2(i_2)), \ldots, \sigma_k^{-1}(\sigma_{k-1}(i_{k-1})) < i_k.
$$

(11)

Since all the components $x_j^i, i \in [n], j \in [d]$, of $X$ are independent random variables, we have that for a fixed index $\nu \in [d]$ each permutation
\[ \tau : [n] \to [n] \] is equally likely to fulfill \( \sigma_j(X) = \tau \). Thus, the probability of \( \nu \) being a \( \delta(X) \)-critical index is just the number of \( k \)-tuples \( (\sigma_1, \ldots, \sigma_k) \) of permutations fulfilling (11), divided by \((n)!^k\).

For given pairwise distinct values \( \sigma_1(i_1), \ldots, \sigma_k(i_k) \) the \( j \)-th condition of (11) is satisfied by \( ((i_j - 1) \ldots (i_j - (k - 1)))(n - k)! \) permutations \( \sigma_j \). Since all \( k \) conditions in (11) can be solved independently of each other, it is now easy to deduce the statement of the Proposition.

To state the corresponding proposition for \( \tilde{\delta} \), we have to introduce \textit{Stirling numbers of second kind} \( S(d, k) \). For \( k \in \mathbb{N}, k \leq d \) let \( S(d, k) \) denote the number of partitions of \([d]\) into \( k \) non-empty subsets. A closed formula for \( S(d, k) \) is

\[
S(d, k) = \sum_{j=0}^{k} \frac{(-1)^j (k - j)^d}{j! (k - j)!}.
\]

This formula and other useful identities can, e.g., be found in [Rio58].

**Proposition 5.5.** Let \( X \) be a uniformly distributed random variable in \([0, 1]^{nd}\). Let \( \nu = (i_1, \ldots, i_d) \in [n]^d \). Then \( \nu \) is a \( \tilde{\delta}(X) \)-critical multi-index with probability

\[
\sum_{k=1}^{d} S(d, k) \left( \frac{(n - k)!}{n!} \right)^{d-1} \prod_{j=1}^{d} \left( \prod_{\nu=1}^{k-1} (i_j - \nu) \right).
\]

**Proof.** We just need to consider the case where the almost-sure event \(|\Gamma_j(X)| = n\) for all \( j \in [d] \) holds. For \( j = 1, \ldots, d \) let \( \sigma_j := \sigma_j(X) : [n] \to [n] \) be the permutation with

\[
x_j^{\sigma_j(1)} < x_j^{\sigma_j(2)} < \cdots < x_j^{\sigma_j(n)} < 1.
\]

Then \( \Phi(\nu)_j = \phi_j(i_j) = x_j^{\sigma_j(i_j)} \) for all \( j \in [d] \). It is easy to see that the surface \( \tilde{S}_j(\Phi(\nu)) \) is \( \tilde{\delta}(X) \)-critical if and only if the condition

\[
\forall j \in [d], \forall l \in [d] \setminus \{j\} : x_l^{\sigma_j(i_l)} \leq \Phi(\nu)_l = x_l^{\sigma_l(i_l)}
\]

is satisfied. This can be rewritten as

\[
\begin{align*}
\sigma_1^{-1}(\sigma_2(i_2)) & , \sigma_1^{-1}(\sigma_3(i_3)) , \ldots , \sigma_1^{-1}(\sigma_d(i_d)) & \leq i_1 , \\
\sigma_2^{-1}(\sigma_1(i_1)) & , \sigma_2^{-1}(\sigma_3(i_3)) , \ldots , \sigma_2^{-1}(\sigma_d(i_d)) & \leq i_2 , \\
\vdots & & \vdots \\
\sigma_d^{-1}(\sigma_1(i_1)) & , \sigma_d^{-1}(\sigma_2(i_2)) , \ldots , \sigma_d^{-1}(\sigma_{d-1}(i_{d-1})) & \leq i_d .
\end{align*}
\]

(12)

If \(|\{\sigma_1(i_1), \ldots, \sigma_d(i_d)\}| = k\), then there exist

\[
S(d, k) n! \left( (n - k)! \right)^{d-1} \prod_{j=1}^{d} \prod_{\nu=1}^{k-1} (i_j - \nu)
\]

25
permutations satisfying (12). With this observation and the fact that all components $x^i_j, i \in [n], j \in [d]$, of $X$ are stochastically independent, it is now easy to deduce the statement of Proposition 5.5.

6 Experimental Results

We now present the experimental evaluations of the algorithms. We will compare our basic and improved algorithms, $TA_{\text{basic}}$ and $TA_{\text{improved}}$, against the algorithm of Winker and Fang [WF97], and also give a brief comparison against the genetic algorithm of Shah [Sha10] and the integer programming-based algorithm of Thiémard [Thi01b].

6.1 Experimental Setup

We divide our experiments into a thorough comparison against the algorithm of Winker and Fang [WF97], given in Section 6.3, and more brief comparisons against the algorithms of Shah [Sha10] and Thiémard [Thi01b], in Sections 6.4 and 6.5, respectively. The algorithms $TA_{\text{basic}}$ and $TA_{\text{improved}}$, as well as the algorithm of Winker and Fang [WF97], were implemented by the authors in the C programming language, based on the code used in [Win07]. All implementations were done with equal care. In the case of Winker and Fang [WF97], while we did have access to the original Fortran source code (thanks to P. Winker), due to lack of compatible libraries we could not use it, and were forced to do a re-implementation.

For the integer programming-based algorithm of Thiémard [Thi01b], E. Thiémard has kindly provided us use of the source code. This source code was modified only as far as necessary for compatibility with newer software versions — specifically, we use version 11 of the CPLEX integer programming package, while the code of Thiémard was written for an older version. Finally, Shah has provided us with the application used in the experiments of [Sha10], but as this application is hard-coded to use certain types of point sets only, we restrict ourselves to comparing with the experimental data published in [Sha10]. Random numbers were generated using the Gnu C library pseudorandom number generator.

The instances used in the experiments are described in Section 6.2. For some instances, we are able to compute the exact discrepancy values either using an implementation of the algorithm of Dobkin et al. [DEM96], available from the third author’s homepage, or via the integer programming-based algorithm of Thiémard [Thi01b]. These algorithms both have far better time dependency than that of Bundschuh and Zhu [BZ93], allowing us to report exact data for larger instances than previously done. For those

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instances where this is too costly, we report instead the largest discrepancy value found by any algorithm in any trial; these imprecise values are marked by a star. Note that this includes some trials with other (more time-consuming) parameter settings than those of our published experiments; thus sometimes, none of the reported algorithms are able to match the approximate max value.

Throughout, for our algorithms and for the Winker and Fang algorithm, we estimate the expected outcome of running 10 independent trials of 100,000 iterations each and returning the largest discrepancy value found, and call this the best-of-10 value. The estimation is computed from a basis of 100 independent trials, as suggested by Johnson [Joh02], which strongly decreases irregularities due to randomness compared to the method of taking 10 independent best-of-10 values and averaging these. The comparisons are based on a fix number of iterations, rather than equal running times, as the point of this paper is to compare the strengths of the involved concepts and ideas, rather than implementation tweaks. For this purpose, using a re-implementation rather than the original algorithm of Winker and Fang [WF97] has the advantage that all algorithms compared use the same code base, compiler, and libraries, including the choice of pseudo-random number generator. This further removes differences that are not interesting to us.

6.2 Instances

Our point sets are of four types: Halton sequences [Hal60], Faure sequences [Fau82], Sobol’ point sets [Sob67], and so-called Good Lattice Points (GLP), described below. The Halton sequences and GLPs were generated by programs written by the authors, the Faure sequences by a program of John Burkardt [Bur], and the Sobol’ sequences using the data and code of Stephen Joe and Frances Kuo [JK08, Kuo10].

Winker and Fang tested their algorithm for several point sets in dimension $d = 4, 5, \ldots, 11$, which were constructed in the following manner: Let $(n, h_1, \ldots, h_d) \in \mathbb{N}^{d+1}$ with $0 < h_1 < h_2 < \cdots < h_d < n$, where at least one $h_i$ is relatively prime with $n$, i.e., their greatest common divisor is one. Then the points $x^1, \ldots, x^n \in [0, 1)^d$ are given by

$$x^i_j := \left\{ \frac{2ih_j - 1}{2n} \right\}, \quad i \in [n], \ j \in [d],$$

where $\{x\}$ denotes the fractional part of $x \in \mathbb{R}$, i.e., $\{x\} = x - \lfloor x \rfloor$. Winker and Fang call $\{x^1, \ldots, x^n\}$ a good lattice point (GLP) set$^4$ of the generating vector $(n, h_1, \ldots, h_d)$. It is known that for any $d \geq 2$ and $n \geq 2$ there exists a generating vector such that the corresponding GLP set exhibits

$^4$Other authors call it GLP set if it additionally exhibits a small discrepancy.
asymptotically a discrepancy of $O(\log(n)^d/n)$, where the implicit constant of the big-O-notation depends solely on $d$, see, e.g., [Nie92, Sect. 5.2].

Winker and Fang considered two series of examples to test their algorithm: First, they (randomly) generated in each dimension $d = 4, 5, 6$ ten GLP $n$-point sets, where $n \in \{50, 51, \ldots, 500\}$ for $d = 4$, $n \in \{50, 51, \ldots, 250\}$ for $d = 5$ and $n \in \{25, 26, \ldots, 100\}$ for $d = 6$. For each GLP set the exact discrepancy was calculated with an implementation of the algorithm of Bundschuh and Zhu [BZ93].

Secondly, they considered six GLP sets in dimension $d = 6, 7, \ldots, 11$ with cardinality between 2129 and 4661 points and performed 20 trials with 200,000 iterations for each of the six sets. Solving these instances exactly is mostly intractable, even with the algorithm of Dobkin et al. [DEM96]. Therefore, with the exception of the smallest instance, it cannot be said if the results of this second series of examples are good approximations of the real discrepancy of the GLP sets under consideration or not.

### 6.3 Comparisons against the Algorithm by Winker and Fang

To begin the comparisons, an indication of the running times of the algorithms is given in Table 1. As can be seen from the table, TA\textsubscript{basic} takes slightly more time than our implementation of Winker and Fang, and TA\textsubscript{improved} takes between two and three times as long, mainly due to the snapping procedures. For TA\textsubscript{improved}, we report the separate times for $\delta$ and $\bar{\delta}$ optimization, as well as the time required for a mixed optimization of both (as is done in TA\textsubscript{basic}). As can be seen, the overhead due to splitting is negligible to non-existent.

The parameter settings for our implementation of the algorithm by Winker and Fang are as follows. Since our experiments did not reveal a strong influence of the choice of $\alpha$ on the quality of the algorithm, we fix $\alpha := 0.995$ for our experiments. Winker and Fang do not explicitly give a rule how one should choose $k$ and $mc$. For the small-dimensional data (Table 2), we use the settings of [WF97]. For the other tests, we use $mc = 3$ if $d \leq 12$ and $mc = 4$ otherwise, and $k = 41$ if $n \leq 500$ and $k = 301$ otherwise. This seems to be in line with the choices of Winker and Fang for the

<table>
<thead>
<tr>
<th>Algorithm</th>
<th>Smaller instance</th>
<th>Larger instance</th>
</tr>
</thead>
<tbody>
<tr>
<td>$d = 10, n = 100$</td>
<td>0.78s</td>
<td>9.34s</td>
</tr>
<tr>
<td>$d = 20, n = 1000$</td>
<td>1.22s</td>
<td>10.94s</td>
</tr>
<tr>
<td>TA\textsubscript{improved}, $\delta$ only</td>
<td>0.85s</td>
<td>9.11s</td>
</tr>
<tr>
<td>TA\textsubscript{improved}, mixed form</td>
<td>1.87s</td>
<td>20.37s</td>
</tr>
<tr>
<td>Winker &amp; Fang</td>
<td>0.61s</td>
<td>7.2s</td>
</tr>
</tbody>
</table>

Table 1: Running times for the considered algorithms. All algorithms executed one trial of 100,000 iterations. The inputs are two randomly generated point sets.
Table 2: Data for GLP sets used by Winker and Fang [WF97]. Discrepancy values marked with a star are lower bounds only (i.e., largest discrepancy found over all executions of algorithm variants). All data is computed using 100 trials of 100,000 iterations; reported is the average value of best-of-10 calls, and number of times (out of 100) that the optimum (or a value matching the largest known value) was found. The data for Winker and Fang is for our re-implementation of the algorithm; the original results for the same instances can be found in [WF97].
sizes used.

Table 2 shows the data for the GLP sets used by Winker and Fang in [WF97]. Although the last group of point sets are quite large, note that this data is mostly of modest dimension. As can be seen, for these sizes, all algorithms behave reasonably, with both of our algorithms generally outperforming our implementation of Winker and Fang, and with TA_{improved} showing much higher precision than TA_{basic}.

We note that our re-implementation of the Winker and Fang algorithm gives notably worse results than what was reported in [WF97] for the same instances. For the larger instances (i.e., with thousands of points), 200,000 iterations are used in [WF97] while we use 100,000 iterations throughout, but there is also a clear difference for the smaller settings. Adjusting the parameters of our implementation to match those used in [WF97] has not been found to compensate for this. After significant experimentation, the best hypothesis we can provide is that there might be a difference in the behavior of the pseudo-random number generators used (in particular, as [WF97] uses a random number library we do not have access to). Still, even compared to the results reported in [WF97], our algorithms, and TA_{improved} in particular, still fare well.

Table 3 shows the new data, for larger-scale instances. A few new trends are noticeable, in particular for the higher-dimensional data. Here, the algorithm of Winker and Fang seems to deteriorate, and there is also a larger difference emerging between TA_{basic} and TA_{improved}, in particular for the Sobol’ sets. However, as can be seen for the 2048-point, 20-dimensional Sobol’ set, it does happen that the lower bound is quite imprecise. (The value of 0.0724 for this point set was discovered only a handful of times over nearly 5000 trials of algorithm variants and settings.)

The highest-dimensional sets (d = 50) illustrate the deterioration of Winker and Fang with increasing dimension; for many of the settings, the largest error this algorithm finds is exactly 1/n (due to the zero-volume box containing the origin with one point).

### 6.4 Comparisons with the Algorithm by Shah

Table 4 lists the point sets used by Shah [Sha10]. The Faure sets here are somewhat nonstandard in that they exclude the origin, i.e., they consist of points 2 through n + 1 of the Faure sequence, where the order of the points is as produced by the program of Burkhardt [Bur].

Some very small point sets were omitted, as every reported algorithm would find the optimum every time. For all but one of the point sets, the exact discrepancy could be computed; the remaining instance is the first 500 points of the 10-dimensional Faure sequence.

Most of the sets seem too easy to really test the algorithms, i.e., all variants frequently find essentially optimal points. The one exception is the
Table 3: New instance comparisons. Discrepancy values marked with a star are lower bounds only (i.e., largest discrepancy found over all executions of algorithm variants). All data is computed using 100 trials of 100,000 iterations; reported is the average value of best-of-10 calls, and number of times (out of 100) that the optimum (or a value matching the largest known value) was found.
Table 4: Comparison against point sets used by Shah. Reporting average value of best-of-10 calls, and number of times (out of 100) that the optimum was found; for Shah, reporting highest value found, and number of times (out of 100) this value was produced. The discrepancy value marked with a star is lower bound only (i.e., largest value found by any algorithm). Values marked (1) are recomputed using the same settings as in [Sha10].

Table 5: Comparison against the integer programming-based algorithm of Thiémard [Thi01b]. The values for \( \text{TA}_{\text{improved}} \) represent the time and average result of a best-of-10 computation with 100,000 iterations per trial. The middle pair of columns give the time required for [Thi01b] to return a first output, and the value of this output; the last two columns report the lower bound reached by [Thi01b] if allocated the same time that \( \text{TA}_{\text{improved}} \) needs for completion, and the time required by [Thi01b] to match the result of \( \text{TA}_{\text{improved}} \). The last item, which shows a clear advantage for our algorithms. We also find (again) that \( \text{TA}_{\text{improved}} \) has a better precision than the other algorithms.

6.5 Comparisons with the Algorithms by Thiémard

Finally, we give a quick comparison against the integer programming-based algorithm of Thiémard [Thi01b]. Since [Thi01b] has the feature that running it for a longer time produces gradually stronger bounds, we report three different checkpoint values; see Table 5 for details. The results are somewhat irregular; however, [Thi01b] may require a lot of time to report a first value, and frequently will not improve significantly on this initial lower bound except after very large amounts of computation time (for example, for the 12-dimensional, 256-point Sobol’s set, the value 0.0872 is discovered in seconds,
while the first real improvement takes over an hour to produce).

Thiémard also constructed a second algorithm for discrepancy estimation, based on delta-covers [Thi01a]; this is freely downloadable from Thiémard’s homepage. Its prime feature is that it provides upper bounds with a non-trivial running time guarantee. The lower bounds that it produces are not as helpful as the upper bounds, e.g., it was reported in [DGW10] and [Sha10] that the lower bounds from the preliminary version of $TA_{\text{basic}}$ [Win07] and the genetic algorithm of Shah [Sha10] are better. Thus we omit this kind of comparison here.

7 Conclusion

Our numerical experiments clearly indicate that the improvements made from the algorithm of Winker and Fang in $TA_{\text{basic}}$ and $TA_{\text{improved}}$ greatly increases the quality of the lower bounds, in particular for the difficult higher-dimensional problem instances. Nevertheless, we do not fully understand the behavior of different algorithm variants with regards to snapping. In particular, one might well have expected the variant described in Section 4.1.1 to do better than the one of Section 4.1 that we currently use. It is still possible that a judicious application of the “snap-move” variant of Section 4.1.1, perhaps only in certain situations, can improve the behavior further.

Still, all in all, we conclude that the algorithms $TA_{\text{basic}}$ and $TA_{\text{improved}}$ presented in the current work represent significant improvements over previous lower-bound heuristics for computing the star discrepancy, and to the best of our knowledge, make up the best performing star discrepancy estimation algorithms available.

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References


A Calculation of $\lambda^d(A_\varepsilon(z))$ and $\pi^d(A_\varepsilon(z))$

Lemma A.1. Let $\varepsilon \in (0,1]$, and let $z \in [0,1]^d$ with $V_z \geq \varepsilon$. Then

$$\lambda^d(A_\varepsilon(z)) = V_z - (V_z - \varepsilon) \sum_{k=0}^{d-1} \frac{(-\ln(1 - \varepsilon/V_z))^k}{k!}.$$  \hfill (13)

Proof. Let $V_z \geq \varepsilon$. Then we have

$$\lambda^d(A_\varepsilon(z)) = \int_{\alpha_1}^{z_1} \int_{\alpha_2}^{z_2} \int_{\alpha_3}^{z_3} \cdots \int_{\alpha_d}^{z_d} d\zeta_d \cdots d\zeta_1,$$

where

$$\alpha_1 = \frac{V_z - \varepsilon}{z_2 z_3 \ldots z_d}, \quad \alpha_2 = \frac{V_z - \varepsilon}{\zeta_1 z_3 \ldots z_d}, \ldots, \quad \alpha_d = \frac{V_z - \varepsilon}{\zeta_1 \zeta_2 \ldots \zeta_{d-1}}.$$

We prove formula (13) by induction over the dimension $d$. If $d = 1$, then clearly $\lambda(A_\varepsilon(z)) = \varepsilon$. Let now $d \geq 2$. We denote by $\tilde{z}$ the $(d-1)$-dimensional vector $(z_2, ..., z_d)$ and by $\tilde{\varepsilon}$ the term $(\varepsilon + (\zeta_1 - z_1)V_{\tilde{z}})/\zeta_1$. Furthermore we define for $i \in [d-1]$ the lower integration limit $\tilde{\alpha}_i = (V_{\tilde{z}} - \tilde{\varepsilon})/(\zeta_2 \cdots \zeta_i + 1 \ldots \zeta_{d-1})$. Note that $\tilde{\alpha}_i = \alpha_{i+1}$. Then, by our induction hypothesis,

$$\lambda^d(A_\varepsilon(z)) = \int_{\alpha_1}^{z_1} \int_{\tilde{\alpha}_1}^{z_1} \int_{\tilde{\alpha}_2}^{z_2} \cdots \int_{\tilde{\alpha}_{d-1}}^{z_{d-1}} d\zeta_d \cdots d\zeta_1 d\zeta_1$$

$$= \int_{\alpha_1}^{z_1} \left( V_\varepsilon - (V_\varepsilon - \tilde{\varepsilon}) \sum_{k=0}^{d-2} \frac{(-\ln(1 - \tilde{\varepsilon}/V_\varepsilon))^k}{k!} \right) d\zeta_1$$

$$= V_\varepsilon - (V_\varepsilon - \varepsilon) - (V_\varepsilon - \varepsilon) \left[ \sum_{k=1}^{d-1} \frac{1}{k!} \ln \left( \frac{V_\varepsilon}{V_\varepsilon - \varepsilon} \right) \right]_{\zeta_1 = \alpha_1}^{z_1}$$

$$= V_\varepsilon - (V_\varepsilon - \varepsilon) \sum_{k=0}^{d-1} \frac{(-\ln(1 - \varepsilon/V_\varepsilon))^k}{k!}.$$

Proposition A.2. Let $d \geq 2$. For $z \in [0,1]^d$ with $V_z > \varepsilon$, we obtain

$$\lambda^d(A_\varepsilon(z)) = \frac{1}{d!} \frac{\varepsilon^d}{V_z^{d-1}} \sum_{k=0}^{\infty} b_k(d) \left( \frac{\varepsilon}{V_z} \right)^k$$

with positive coefficients

$$b_k(2) = \frac{2}{(k+1)(k+2)}, \quad b_k(3) = \frac{6}{(k+2)(k+3)} \sum_{\nu=0}^{k} \frac{1}{\nu+1}.$$
and
\[ b_k(d) = \frac{d!}{(k + d - 1)(k + d)} \sum_{k_1=0}^{k} \cdots \sum_{k_{d-2}=0}^{k_{d-3}} \prod_{j=1}^{d-2} \frac{1}{k_j + d - j - 1} \] for \( d \geq 4 \).

The power series converges for each \( \epsilon > 0 \) uniformly and absolutely for all \( V_z \in [\epsilon + \epsilon, 1] \). Furthermore, we have \( b_0(d) = 1 \), \( b_1(d) = d(d-1)/2(d+1) \), and for all \( k \) the inequality \( b_k(d) \leq d^k/2^{k-1} \) is satisfied.

**Proof.** To prove the power series expansion, we consider the function
\[ R(x, d) = 1 - (1-x) \sum_{k=0}^{d-1} \frac{(-\ln(1-x))^k}{k!} \] for \( x \in [0,1) \).

Due to Lemma A.1 we have \( \chi^d(A_z(z)) = V_z R(\epsilon/V_z, d) \). Since \( \partial_x R(x, d) = (-\ln(1-x))^{d-1}/(d-1)! \), it suffices to prove the following statement by induction over \( d \):
\[ (-\ln(1-x))^{d-1} = \frac{1}{d!} \sum_{k=0}^{\infty} (k+d)b_k(d)x^{k+d-1}, \quad (14) \]
where the power series converges for each \( \epsilon > 0 \) uniformly and absolutely on \([0, 1 - \epsilon]\). Let first \( d = 2 \). Then
\[-\ln(1-x) = \sum_{k=1}^{\infty} \frac{x^k}{k} = \frac{1}{2!} \sum_{k=0}^{\infty} (k+2)b_k(2)x^{k+1}, \]
and the required convergence of the power series is obviously given. Now let \( d \geq 3 \). Our induction hypothesis yields
\[ \partial_x \frac{(-\ln(1-x))^{d-1}}{(d-1)!} = \frac{1}{1-x} \frac{(-\ln(1-x))^{d-2}}{(d-2)!} \]
\[ = \left( \sum_{\mu=0}^{\infty} x^\mu \right) \left( \frac{1}{(d-1)!} \sum_{\mu=0}^{\infty} (\mu + d - 1)b_\mu(d-1)x^{\mu+d-2} \right) \]
\[ = \frac{1}{d!} \sum_{k=0}^{\infty} \left( d \sum_{\mu=0}^{k} (\mu + d - 1)b_\mu(d-1) \right) x^{k+d-2}, \]
where the last power series converges as claimed above. Now
\[ d \sum_{\mu=0}^{k} (\mu + d - 1)b_\mu(d-1) = \sum_{\mu=0}^{k} \frac{d!}{(\mu + d - 2)} \sum_{\mu_1=0}^{\mu} \cdots \sum_{\mu_{d-3}=0}^{\mu_{d-4}} \prod_{j=2}^{d-3} \frac{1}{\mu_j + d - 2 - j} \]
\[ = d! \sum_{\nu_1=0}^{k} \sum_{\nu_2=0}^{\nu_1} \cdots \sum_{\nu_{d-3}=0}^{\nu_{d-2}} \prod_{j=1}^{d-2} \frac{1}{\nu_j + d - j - 1} \]
\[ = (k+d)(k+d-1)b_k(d). \]
After integration we get (14).

Furthermore, it is easily seen that $b_0(d) = 1$ and $b_1(d) = d(d-1)/2(d+1)$. To complete the proof, we verify $b_k(d) \leq d^k/2^{k-1}$ for $k \geq 2$. The inequality is obviously true in dimension $d = 2$. Hence let $d \geq 3$. From the identity

$$
\sum_{\nu_1=0}^{k} \ldots \sum_{\nu_{d-2}=0}^{d-3} \frac{1}{\nu_j + d - 1 - j} \leq \frac{1}{(d-2)!} \left( \frac{k + d - 2}{k} \right),
$$

we obtain

$$
\sum_{\nu_1=0}^{k} \sum_{\nu_2=0}^{d-3} \ldots \sum_{\nu_{d-2}=0}^{d-2} \frac{1}{\nu_j + d - 1 - j} \leq \frac{1}{(d-2)!} \left( \frac{k + d - 2}{k} \right),
$$

which leads to

$$
b_k(d) \leq \frac{d(d-1)}{(k+d)(k+d-1)} \frac{(k+d-2)...(1+d-2)}{k...1} \leq \frac{d(d-1)}{(k+d)(k+d-1)} \left( \frac{d}{2} \right)^{k-1} (d-1) \leq \frac{d^k}{2^{k-1}}.\]

\[\square\]

Corollary A.3. Let $z \in [0, 1]^d$. If $V_z \geq d \varepsilon$, then

$$
\lambda^d(A_{\varepsilon}(z)) \leq \frac{5}{2d!} \frac{\varepsilon^d}{V_z^{d-1}}.
$$

We now consider the polynomial product measure $\pi^d$.

Lemma A.4. Let $\varepsilon \in (0, 1]$, and let $z \in [0, 1]^d$ with $V_z \geq \varepsilon$. Then

$$
\pi^d(A_{\varepsilon}(z)) = V_z^d - (V_z - \varepsilon)^d \sum_{k=0}^{d-1} \frac{d^k}{k!} (-\ln(1 - \varepsilon/V_z))^k,
$$

and, as a function of $V_z$, $\pi^d(A_{\varepsilon}(z))$ is strictly increasing.

Proof. Let $V_z \geq \varepsilon$. We have

$$
\pi^d(A_{\varepsilon}(z)) = \int_{A_{\varepsilon}(z)} d^dV_z^{d-1} \lambda^d(dx) = \int_0^\varepsilon G(V_z, r) \, dr,
$$

where

$$
G(V_z, r) := d^d(V_z - r)^{d-1} \partial_r \lambda^d(A_r(z)).
$$

From (13) we get for all $0 \leq r \leq \varepsilon$

$$
\partial_r \lambda^d(A_r(z)) = \frac{(-\ln(1 - r/V_z))^{d-1}}{(d-1)!}.
$$

40
If we define
\[ F(r) := -(Vz - r)^d \sum_{k=0}^{d-1} \frac{d^k}{k!} (-\ln(1 - r/Vz))^k, \]
then we observe that \( F'(r) = G(Vz, r) \) holds. Thus we have \( \pi^d(A_\varepsilon(z)) = F(\varepsilon) - F(0) \), which proves (16). Furthermore, according to (17), we get
\[ \partial \nu_\varepsilon \pi^d(A_\varepsilon(z)) = \int_0^\varepsilon \partial \nu_\varepsilon G(Vz, r) \, dr. \]
The integrand of the integral is positive, as the next calculation reveals:
\[
\partial \nu_\varepsilon G(Vz, r) = d^d(Vz - r)^{d-2} \frac{(-\ln(1 - r/Vz))^{d-2}}{(d - 2)!} (-\ln(1 - r/Vz) - r/Vz)
\]
\[ = d^d(Vz - r)^{d-2} \frac{(-\ln(1 - r/Vz))^{d-2}}{(d - 2)!} \sum_{k=2}^{\infty} \frac{1}{k} \left( \frac{r}{Vz} \right)^k > 0 \]
for all \( 0 < r < Vz \). Thus \( \partial \nu_\varepsilon \pi^d(A_\varepsilon(z)) > 0 \), and, considered as a function of \( Vz \), \( \pi^d(A_\varepsilon(z)) \) is strictly increasing.

**Proposition A.5.** Let \( \varepsilon \in (0, 1) \), and let \( z \in [0, 1]^d \) with \( Vz \geq \varepsilon \). Then we have the lower bound \( \pi^d(A_\varepsilon(z)) \geq \varepsilon^d \). If furthermore \( Vz \geq d \varepsilon \), then we have the estimate
\[
e^{-1} \frac{d^d}{d!} \varepsilon^d \leq \pi^d(A_\varepsilon(z)) \leq \frac{5d^d}{2d!} \varepsilon^d.
\]
**Proof.** Let \( Vz = \varepsilon \). Then
\[ \pi^d(A_\varepsilon(z)) = \int_{[0, z]} d^dV_x^{d-1} \lambda^d(dx) = \prod_{i=1}^{d} z_i^d = \varepsilon^d. \]
Since \( \pi^d(A_\varepsilon(z)) \) is an increasing function of \( Vz \), the first lower bound holds. Let now \( Vz \geq d \varepsilon \). Here we use the simple estimate
\[ d^d(Vz - \varepsilon)^{d-1} \lambda^d(A_\varepsilon(z)) \leq \pi^d(A_\varepsilon(z)) \leq d^dVz^{d-1} \lambda^d(A_\varepsilon(z)). \]
Together with Proposition A.2 and Corollary A.3 this leads to
\[
e^{-1} \frac{d^d}{d!} \varepsilon^d \leq (1 - 1/d)^{d-1} \frac{d^d}{d!} \varepsilon^d \leq \pi^d(A_\varepsilon(z)) \leq \frac{5d^d}{2d!} \varepsilon^d. \]

**Remark A.6.** Like \( \lambda^d(A_\varepsilon(z)) \) in Proposition A.2, one can also expand \( \pi^d(A_\varepsilon(z)) \) into a power series. This leads to
\[ \pi^d(A_\varepsilon(z)) = \frac{d^d}{d!} \varepsilon^d \sum_{k=0}^{\infty} a_k(d) \left( \frac{\varepsilon}{Vz} \right)^k, \]
but here the coefficients \( a_k(d) \) are not all positive. So we have, e.g., \( a_0(d) = 1 \), but \( a_1(d) = -d(d - 1)/2(d + 1) \). Therefore the power series expansion is here less useful than in the situation of Proposition A.2.