Algorithms are integral to computer science and every computer scientist (even as an undergrad) has designed several algorithms. So has many a physicist, electrical engineer, mathematician etc. This course is meant to be your one-stop shop that teaches you how to design a variety of algorithms. The operative word is “variety.” In other words you will avoid the blinders that one often sees in domain experts. A bayesian needs priors on the data before he can design algorithms; an optimization expert wishes to cast all problems as mathematican optimization; a systems designer has never seen any problem that cannot be solved by hashing. (OK, mostly kidding but the joke does reflect truth to some degree.) These and more domain-specific ideas make an appearance in our course, but we will learn to not be wedded to any single approach.

The primary skill you will learn in this course is how to analyze algorithms: prove their correctness and their running time and any other relevant properties. Learning to analyze a variety of algorithms (designed by others) will let you design better algorithms later in life. I will try to fill the course with beautiful algorithms. Be prepared for frequent rose-smelling stops, in other words.

1 Difference between grad and undergrad algorithms

Undergrad algorithms is largely about algorithms discovered before 1990; grad algorithms is a lot about algorithms discovered since 1990. OK, I picked 1990 as an arbitrary cutoff. Maybe it is 1985, or 1995. What happened in 1990 that caused this change, you may ask? Nothing. It was no single event but just a gradual shift in the emphasis and goals of computer science as it became a more mature field.

In the first few decades of computer science, algorithms research was driven by the goal of understanding how to design basic components of a computer: operating systems, compilers, networks, etc. Other motivations to study algorithms came out of discrete mathematics, operations research, graph theory. Thus in undergraduate algorithms you would study data structures, graph traversal, string matching, parsing, network flows, etc. Starting around 1990 theoretical computer science broadened its horizons and started looking at new problems: algorithms for bioinformatics, algorithms and mechanism design for e-commerce, algorithms to understand big data or big networks. This changed algorithms research and the change is ongoing. One big change is that it is often unclear what the algorithmic problem even is. Identifying it is part of the challenge. Thus good modeling is important. This in turn is shaped by understanding what is possible (given our understanding of computational complexity) and what is reasonable given the limitations of the type of inputs we are given.
Some examples of this change:

The changing graph. In undergrad algorithms the graph is given and arbitrary (worst-case). In grad algorithms we are willing to look at where the graph came from (social network, computer vision etc.) since those properties may be germane to designing a good algorithm. (This is not a radical idea of course but we will see that formulating good graph models is not easy. This is why you see a lot of heuristic work in practice, without any mathematical proofs of correctness.)

Changing data structures: In undergrad algorithms the data structures were simple and often designed to hold data generated by other algorithms. A stack allows you to hold vertices during depth-first search traversal of a graph, or instances of a recursive call to a procedure. A heap is useful for sorting and searching.

But in the newer applications, data often comes from sources we don’t control. Thus it may be noisy, or inexact, or both. It may be high dimensional. Thus something like heaps will not work, and we need more advanced data structures.

We will encounter the “curse of dimensionality” which constrains algorithm design for high-dimensional data.

Changing notion of input/output: Algorithms in your undergrad course have a simple input/output model. But increasingly we see a more nuanced interpretation of what the input is: datastreams (useful in analytics involving routers and web servers), online (sequence of requests), social network graphs, etc. And there is a corresponding subtlety in settling on what an appropriate output is, since we have to balance output quality with algorithmic efficiency. In fact, design of a suitable algorithm often goes hand in hand with understanding what kind of output is reasonable to hope for.

Type of analysis: In undergrad algorithms the algorithms were often exact and work on all (i.e., worst-case) inputs. In grad algorithms we are willing to relax these requirements.
1 Preliminaries

In hashing, we want to store a subset $S$ of a large universe $U$ ($U$ can be very large, say $|U| = 2^{32}$ is the set of all 32 bit integers). And $|S| = m$ is a relatively small subset. For each $x \in U$, we want to support 3 operations:

- **insert**($x$). Insert $x$ into $S$.
- **delete**($x$). Delete $x$ from $S$.
- **query**($x$). Check whether $x \in S$.

A hash table can support all these 3 operations. We design a hash function

$$h : U \rightarrow \{0, 1, \ldots, n - 1\}$$

(1.1)

such that $x \in U$ is placed in $T[h(x)]$, where $T$ is a table of size $n$.

Since $|U| \gg n$, multiple elements can be mapped into the same location in $T$, and we deal with these collisions by constructing a linked list at each location in the table.

One natural question to ask is: how long is the linked list at each location? We make two kinds of assumptions:
1. Assume the input is the random.

2. Assume the input is arbitrary, but the hash function is random.

Assumption 1 may not be valid for many applications, since the input might be correlated.

For Assumption 2, we construct a set of hash functions $\mathcal{H}$, and for each input, we choose a random function $h \in \mathcal{H}$ and hope that on average we will achieve good performance.

## 2 Hash Functions

Say we have a family of hash functions $\mathcal{H}$, and for each $h \in \mathcal{H}$, $h : U \rightarrow [n]^1$, what do mean by saying these functions are random?

For any $x_1, x_2, \ldots, x_m \in S$ ($x_i \neq x_j$ when $i \neq j$), and any $a_1, a_2, \ldots, a_m \in [n]$, ideally a random $\mathcal{H}$ should satisfy:

- $\Pr_{h \in \mathcal{H}}[h(x_1) = a_1] = \frac{1}{n}$.
- $\Pr_{h \in \mathcal{H}}[h(x_1) = a_1 \land h(x_2) = a_2] = \frac{1}{n^2}$. Pairwise independence.
- $\Pr_{h \in \mathcal{H}}[h(x_1) = a_1 \land h(x_2) = a_2 \land \cdots \land h(x_k) = a_k] = \frac{1}{n^k}$. $k$-wise independence.
- $\Pr_{h \in \mathcal{H}}[h(x_1) = a_1 \land h(x_2) = a_2 \land \cdots \land h(x_m) = a_m] = \frac{1}{n^m}$. Full independence (note that $|U| = m$). In this case we have $n^m$ possible $h$ (we store $h(x)$ for each $x \in U$), so we need $m \log n$ bits to represent the each hash function. Since $m$ is usually very large, this is not practical.

For any $x$, let $L_x$ be the length of the linked list containing $x$, then $L_x$ is just the number of elements with the same hash value as $x$. Let random variable

$$I_y = \begin{cases} 
1 & \text{if } h(y) = h(x), \\
0 & \text{otherwise.} 
\end{cases} \quad (2.1)$$

So $L_x = 1 + \sum_{y \neq x} I_y$, and

$$E[L_x] = 1 + \sum_{y \neq x} E[I_y] = 1 + \frac{m-1}{n} \quad (2.2)$$

Note that we don’t need full independence to prove this property, and pairwise independence would actually suffice.

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1We use $[n]$ to denote the set $\{0, 1, \ldots, n-1\}$
3 2-Universal Hash Families

**Definition 3.1** (Cater Wegman). Family $\mathcal{H}$ of hash functions is 2-universal if for any $x \neq y \in U$,
\[
\Pr_{h \in \mathcal{H}}[h(x) = h(y)] \leq \frac{1}{n} \tag{3.1}
\]

Note that this property is even weaker than 2 independence.

We can design 2-universal hash families in the following way. Choose a prime $p \in \{\lvert U \rvert, \ldots, 2\lvert U \rvert\}$, and let
\[
f_{a,b}(x) = ax + b \mod p \quad (a, b \in [p], a \neq 0) \tag{3.2}
\]
And let
\[
h_{a,b}(x) = f_{a,b}(x) \mod n \tag{3.3}
\]

**Lemma 3.2.** For any $x_1 \neq x_2$ and $s \neq t$, the following system
\[
\begin{align*}
ax_1 + b &= s \mod p \tag{3.4} \\
ax_2 + b &= t \mod p \tag{3.5}
\end{align*}
\]
has exactly one solution.

Since $[p]$ constitutes a finite field, we have that $a = (x_1 - x_2)^{-1}(s - t)$ and $b = s - ax_1$. Since we have $p(p-1)$ different hash functions in $\mathcal{H}$ in this case,
\[
\Pr_{h \in \mathcal{H}}[h(x_1) = s \land h(x_2) = t] = \frac{1}{p(p-1)} \tag{3.6}
\]

**Claim 3.3.** $\mathcal{H} = \{h_{a,b} : a, b \in [p] \land a \neq 0\}$ is 2-universal.

**Proof.** For any $x_1 \neq x_2$,
\[
\begin{align*}
\Pr[h_{a,b}(x_1) = h_{a,b}(x_2)] &= \sum_{s, t \in [p], s \neq t} \delta_{s = t \mod n} \Pr[f_{a,b}(x_1) = s \land f_{a,b}(x_2) = t] \tag{3.7} \\
&= \frac{1}{p(p-1)} \sum_{s, t \in [p], s \neq t} \delta_{s = t \mod n} \tag{3.8} \\
&\leq \frac{1}{p(p-1)} \frac{p(p-1)}{n} \tag{3.9} \\
&= \frac{1}{n} \tag{3.10}
\end{align*}
\]

where $\delta$ is the Dirac delta function. Equation (3.10) follows because for each $s \in [p]$, we have at most $(p-1)/n$ different $t$ such that $s \neq t$ and $s = t \mod n$. \qed
Can we design a collision free hash table then? Say we have $m$ elements, and the hash table is of size $n$. Since for any $x_1 \neq x_2$, $\Pr[h(x_1) = h(x_2)] \leq \frac{1}{n}$, the expected number of total collisions is just

$$E[\sum_{x_1 \neq x_2} h(x_1) = h(x_2)] = \sum_{x_1 \neq x_2} E[h(x_1) = h(x_2)] \leq \binom{m}{2} \frac{1}{n} \quad (3.12)$$

Let's pick $m \geq n^2$, then

$$E[\text{number of collisions}] \leq \frac{1}{2} \quad (3.13)$$

and so

$$\Pr_{h \in H}[\exists \text{ a collision}] \leq \frac{1}{2} \quad (3.14)$$

So if the size the hash table is large enough $m \geq n^2$, we can easily find a collision free hash functions. But in reality, such a large table is often unrealistic. We may use a two-layer hash table to avoid this problem.

![Figure 2: Two layer hash tables.](image)

Specifically, let $s_i$ denote the number of collisions at location $i$. If we can construct a second layer table of size $s_i^2$, we can easily find a collision-free hash table to store all the $s_i$ elements. Thus the total size of the second-layer hash tables is $\sum_{i=0}^{m-1} s_i^2$.

Note that $\sum_{i=0}^{m-1} s_i(s_i - 1)$ is just the number of collisions calculated in Equation (3.12), so

$$E[\sum_{i} s_i^2] = E[\sum_{i} s_i(s_i - 1)] + E[\sum_{i} s_i] = \frac{m(m-1)}{n} + m \leq 2m \quad (3.15)$$

4 Load Balance

In load balance problem, we can imagine that we are trying to put balls into bins. If we have $n$ balls and $n$ bins, and we randomly put the balls into bins,
then for a give \( i \),

\[
\Pr[\text{bin}_i \text{ gets more than } k \text{ elements}] \leq \binom{n}{k} \cdot \frac{1}{n^k} \leq \frac{1}{k!} \quad (4.1)
\]

By Stirling’s formula,

\[
k! \sim \sqrt{2\pi n} \left( \frac{k}{e} \right)^k \quad (4.2)
\]

If we choose \( k = O\left( \frac{\log n}{\log \log n} \right) \), we can let \( \frac{1}{k^2} \leq \frac{1}{n^2} \). Then

\[
\Pr[\exists \text{ a bin } \geq k \text{ balls}] \leq n \cdot \frac{1}{n^2} = \frac{1}{n} \quad (4.3)
\]

So with probability larger than \( 1 - \frac{1}{n^2} \),

\[
\max \text{ load} \leq O\left( \frac{\log n}{\log \log n} \right) \quad (4.4)
\]

Note that if we look at 2 random bins when a new ball comes in and put the ball in the bin with fewer balls, we can achieve maximal load at the scale of \( O(\log \log n) \), which is a huge improvement.

\[\text{This can be easily improve to } 1 - \frac{1}{n^c} \text{ for any constant } c\]