Introduction to Stochastic Process STAT 615 Course Notes

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0 Preliminaries

Definition. 0.1. A <u>sample space</u> Ω is the collection of individual <u>outcomes</u> or <u>realizations</u> for a random experiment.

Subsets of Ω are called <u>events</u>. For $A \subset \Omega$, we say "A occurs" if the outcome $\omega \in A$

 Ω can be <u>countable</u> (finite or countably infinite) or <u>uncountable</u>. It need not consist of real-valued entities. Most often, its role is in the background and simply assured.

Example. 0.1. Toss a coin infinitely many times, observing 1 (heads) or 0 (tails) for each toss. The outcomes are inifinite sequences of 0's & 1's, such as $(1, 1, 0, 0, 0, 1, \cdots)$. The collection Ω of all possible outcomes is uncountable in this case.

Events are subsets of Ω , often identified descriptively:

• "1st toss is H"= $\{(x_1, x_2, \cdots) : x_1 = 1\}$.

- "12 H's in the 1st 100 tosses" = $\{(x_1, x_2, \dots) : \sum_{i=1}^{1} 00x_i = 12\}$.
- "the long tem proportion of H's is $\frac{1}{3}$ " = $\{(x_1, x_2, \cdots) : \frac{1}{n} \sum_{i=1}^n x_i \to \frac{1}{3}\}.$

A <u>singleton</u> event is a subset of just one outcome, which is not quite the same as the outcome itself.

The empty set ϕ is considered to be an event also.

Definition. 0.2. A <u>probability measure</u> P is a set function (i.e. P is applied to events, not to outcomes) such that

(i)
$$0 \le P(A) \le 1$$
, with $P(\Omega) = 1$, $P(\phi) = 0$

(ii) if A_1, A_2, \cdots are <u>disjoint</u> then $P(\bigcup_{i=1}^{\infty} A_i) = \sum_{i=1}^{\infty} P(A_i)$ (similarly for finite unions) when the A_i 's are not disjoint, we have

Theorem 0.1. (Boole's inequality)

$$P(\bigcup_{i=1}^{n} A_i) \le \sum_{i=1}^{n} P(A_i)$$

.

Definition. 0.3. A collection of **A** of subsets of Ω is called a σ -algebra (or σ -field) if

- (i) $\phi, \Omega \in \mathbf{A}$
- (ii) $A \in \mathcal{A} \Longrightarrow A^{c} \in \mathcal{A}$
- (iii) $A_1, A_2, \dots \in \mathcal{A} \Longrightarrow \bigcup_{i=1}^{\infty} A_i \in \mathcal{A}$ (Some for finite unions) and hence also $\bigcap_{i=1}^{\infty} \in \mathcal{A}$

Note: the requirement is closure under <u>countable</u> unions and intersections. So a union of uncountably may subsets in \mathcal{A} need not itself be in \mathcal{A} . (there are important mathematical reasons for allowing this.) **Additional Note:** <u>complement</u> A^{\downarrow} ="A does not occur" union of events $\bigcup_i A_i$ = "some A_i (at least one) occurs" intersection $\bigcap_i A_i$ = "every A_i occurs"

Example. 0.2. If Ω is countable (such as the integers \mathbb{Z}) then we can let \mathcal{A} be all subsets.

Example. 0.3. If $\Omega = \mathbb{R}$ (all real values), we usually use the <u>Borel Sets</u> $B = \text{smallest } \sigma$ -algebra that contains all singletons and all intervals. So \mathcal{B} would include countable unions of intervals, but this by no means comes close to all it has. There is no direct formula or description to characterize all arbitrary Borel sets.

On the other hand, given $B \in \mathcal{B}$ and a probability measure P, P(B) can be approximated as close as we like with the probability of some finite union of intervals.

Borel sets on \mathbb{R}^d (d-dimensional vectors) are defined similarly, using rectangles.

Example. 0.4. (cond.) Let $\Omega = \{(x_1, x_2, x_3, \cdots) : x_i = 0 \text{ or } x_i = 1 \text{ for each } i \geq 1\}$ be the sample space of infinite sequences of coin tosses by <u>cylinder events</u> (events where a <u>finite</u> number of the x_i 's are fixed). For example,

$$\{(x_1, x_2, \cdots) : x_1 = 1, x_2 = 0\}$$

.

The triple (Ω, \mathcal{A}, P) (i.e. sample space, σ -algebra of events probability measure) is called a probability space.

Definition. 0.4. A <u>random variable</u> $X = X(\omega)$ is a real-valued function applied to outcomes (i.e., $X : \Omega \to \mathbb{R}$.

An <u>extended random variable</u> can possibly take infinite values (i.e., $X: \Omega \to \mathbb{R} \stackrel{def}{=} \mathbb{R} \lor \{-\infty,\infty\}$ (" $\stackrel{def}{=}$ " means "defined as", the same as "=:"))

Since random variables (rvs) are functions, it is <u>highly recommended</u> that you take care to distinguish them from actual (possible) values. We will usually, but not always, use upper case for rvs and lower case for actual values. (Also, try to avoid using the same name, such as X or Y, for everything, and centatinly make distinctions between rvs in the same problem/context.)

A <u>random vector</u> is a vector of rvs and a <u>random (stochastic) process</u> is a sequence of rvs (e.g. (x_1, x_2, \cdots)) or a function that takes random values (e.g. $x(t), t \in \mathbb{R}$, where each X(t) is a random variable).

Definition. 0.5. Every rv X has a <u>cululative distribution function</u> (cdf)

$$F_X(x) = P(X \le x)$$

(sensible even for extended rvs)

A rv X (or its cdf) is <u>discrete</u> if there is a countable set of values $\{a_i\}$ such that $\sum_i P(x=a_i) = 1, \{P_i\} = \{P(x=a_i)\}$ is called the <u>probability mass function</u> (pmf).

A rv X (or its cdf) is absolutely continuous if

$$P(a < X \le b) = F(b) - F(a) = \int_a^b f(x)dx$$

for some function f(x), called the probability density function (pdf)

<u>However</u>, we do not with to limit ourselves to just two types. There are, for example, rvs with neither a pmf or a pdf, and these are mixtures of types. Things become more general still for random vectors.

Furthermore, a stochastic process cannot be characterized so simply at all (More on this in the 1st chapter).

We will write $X \sim F_X$ (or $X \sim \{P_i\}$ or $X \sim f(x)$ when the context is clear) to signify that X has distribution given by F_x .

You should be familiar with all the commonly used distributions.

• ex. $Y \sim Poisson(\lambda)$ with pmf

$$P(y = k) = \frac{\lambda^k e^{-\lambda}}{k!}, k = 0, 1, 2, \dots$$

.

• ex. $T \sim Gamma(\alpha, \beta)$ with pdf

$$f(t) = \frac{\beta^{\alpha} t^{\alpha - 1} e^{-\beta t}}{\Gamma(\alpha)}, t \ge 0$$

.

Definition. 0.6. Random variables X_1, \dots, X_m are independent if

$$P(X_1 \in A_1, X_2 \in A_2, \dots, X_m \in A_m) = \prod_{i=1}^m P(X_i \in A)$$

for all **Borel sets** A_1, \dots, A_m .

Equivalently,

$$P(X_1 \le x_1, \dots, X_m \le x_m) = \prod_{i=1}^m P(X_i \le x_i)$$

for all real values x_1, \dots, x_m .

(The 1st definition can be applied to random elements of any kind, with appropriate A_i 's.)

A collection $\{X_t\}$ of rvs (possibly uncountably many) is <u>independent</u> if every finite sub-collection is. That is, if X_{t_1}, \dots, X_{t_m} are independent, for any t_1, \dots, t_m .

A sequence $\{X_n\}$ is independent and identically distributed (iid) if it is independent and all the rvs have the same distribution. We sometimes write $\{X_n\} \stackrel{iid}{\sim} F$ (or just $X_n \stackrel{iid}{\sim} F$).

Stochastic processes are generally <u>not independent</u> (they wouldn't be interesting if they were). But they often can be constructed or described in terms of independent rvs.

Random variables and their distributions can also be characterized in terms of expectations. As before, we do not wish to be limited to special types (and in fact it is notationally more convenient not to be).

Definition. 0.7. Here is how the expectation E(g(x)) is defined (if possible).

For simplicity, just let Y = g(X) and we'll define E(Y).

1. indicator rv

if

$$Y = \mathbb{1}_A = \begin{cases} 0, & \text{if A does not occur} \\ 1, & \text{if A does occur} \end{cases}$$

then

$$E(\mathbb{1}_A) = P(A)$$

. indicator rvs have Bernoulli distribution (which simply means discrete on $\{0,1\}$)

2. simple rv

Y is a finite linear combination of indicator rvs

i.e. $Y = \sum_{i=1}^m c_i \mathbb{1}_{A_i}$ (hence discrete w/ finitely many values)

then

$$E(Y) = \sum_{i=1}^{m} c_i P(A_i)$$

. (this works even if the A_i 's are not disjoint or if the c_i 's are not distinct.)

3. nonnegative rv

 $Y \geq 0$. There always exist nonnegative simple rvs $Y_n(\omega) \uparrow Y(\omega)$ for every $\omega \in \Omega$.

Then

$$E(Y) = \lim_{n \to \infty} E(Y_n)$$
 (which myght equal ∞)

(This does not depend on the choice of sequenc $\{Y_n\}$, nor is it necessarily the best way to compute E(y).)

4. general rv Y

Define the positive part of Y, $Y_+ = \max(Y, 0)$

and the negative part, $Y_{-} = \max(-Y, 0)$.

(both are nonnegative)

so
$$Y=Y_+-Y_-$$
 and $|Y|=Y_++Y_-$

Then if either (or both) of $E(Y_+)\&E(Y_-)$ are finite, define

$$E(Y) = E(Y_{+}) - E(Y_{-})$$

. (Again, the value can be infite - we just don't allow the case $\infty - \infty$)

The purpose of the above is just so we can define (and talk about) expectations without having to worry about the type of rv or how the expectation is to be computed. In fact, computing expectations will depend on context and often is not necessary in specific cases.

Definition. 0.8. (Lebesque-Stielties notation).

If $X \sim F$, it is conventional to express

$$E(x) = \int_{-\infty}^{\infty} x F(dx)$$

, and $E(g(x))=\int_{-\infty}^{\infty}g(x)F(dx),$ when they can be defined.

A similar usage applies to random vectors, using a joint cdf (but not to stochastic processes) .

Other terminology.

- A random variable X is finite if $|X(\omega)| < \infty$ for all $\omega \in \Omega$.
- A random variable X is <u>bounded</u> if there exists a constant C such that $|X(\omega)| \leq C$ for all $\omega \in \Omega$.
- Do not confuse "finite" & "bounded".

The terms apply in similar fashion to any real valued function g(x).

An event A occurs almost surely (a.s.) or with probability 1 (w.p.1) if P(A) = 1.

Note that "almost sure" is in the context of a specific probability measure or model, whereas the definition of A (as some event) is not.

ex. Y is almost surely finite if $P(|Y| < \infty) = 1$. However, the event $\{Y = \infty\}$ need not be empty. Indeed, we will want to discuss rvs taht are almost surely finite for some models but not for others (and to identify the distinguishing condition).

ex. X_n converges to X almost surely $(X_n \to X \text{ a.s.})$, if $P(\lim_{n \to \infty} X_n = X) = 1$,

This is stronger than convergence in probability: $P(|X_n - X| > \epsilon) \to 0$ for all positive ϵ .

The basic limit theorems

Theorem 0.2. Monotone Convergence (MCT)

For $0 \le X_n \uparrow X : E(X_n) \to E(X)$ (even if infinite)

Likewise, if $0 \le g_n(x) \uparrow g(x)$ for all x, then

$$\int g_n(x)dx \to \int g(x)dx$$

, and

$$\sum_{i=1}^{\infty} g_n(i) \to \sum_{i=1}^{\infty} g(i)$$

Theorem 0.3. *Dominated Convergence* (DCT)

For $X_n \to X$ such that $|X_n| \le Y$ and $E(y) < \infty$, $E(X_n) \to E(X)$.

(again, similar statements for integrals & sums)

Theorem 0.4. <u>Fatou's Lemma</u> $X_n \ge 0, X_n \to X$

$$\liminf_{n \to \infty} E(X_n) \ge E(X)$$

(& similarly for sums and integrals)

(Note:

$$\liminf_{n \to a_n} = \lim_{n \to \infty} (\inf_{k \ge n} a_k) \quad \& \quad \limsup_{n \to \infty} a_n = \lim_{n \to \infty} (\sup_{k \ge n} a_k)$$

.)

Theorem 0.5. Strong Law of Large Numbers (SLLN) For iid X_n ,

$$\frac{1}{n} \sum_{i=1}^{n} X_i \to E(X) \qquad (if defined)$$

almost surely, and

$$E(|\frac{1}{n}\sum_{i=1}nX_i - E(X)|) \to 0 \quad (if E(|x|) < \infty)$$

.

Equalities & inequalities

Theorem 0.6.

- If $X \ge 0$ then $E(X) = \int_0^\infty P(X > x) dx$, even if infinite (& valid even if $P(X = \infty) > 0$).
- If $X \ge 0$ and is integer-valued then

$$E(X) = \sum_{n=0}^{\infty} P(X > n)$$

Theorem 0.7.

(i) (Markov) If $g(x) \ge 0$ & nondecreasing then

$$P(X \geq x) \leq \frac{E(g(X))}{g(x)}$$

e.g.

$$P(|X| > x) \le \frac{E(|x|)}{x}$$
 & $P(|X| > x) \le \frac{E(x^2)}{x^2}$

.

- (ii) $|E(x)| \le E(|x|)$
- (iii) (Lyapunov) $E(|X|^r) \le (E(x^2)E(y^2))^{\frac{1}{2}}$, if 0 < r < s.
- (iv) (Cauchy-Schwaiz) $|E(xy)| \le (E(x^2)(y^2))^{\frac{1}{2}}$
- (v) (Jenson) If g(x) is convex then $E(g(x)) \ge g(E(x))$ with equality only if g(x) = a + bx for some constants a, b.

Note: "convex" means $g(ax + (1 - a)y) \le ag(x) + (1 - a)g(y), 0 < a < 1$

Equivalently, $g(b) - g(a) = \int_a^b g'(u) du$ all a, b and g'(u) is nondecreasing (the derivative need not exist at all values of u for this to hold)

Theorem 0.8. (Kolmogorov maximal inequality)

If X_n are independent, each $E(X_n^2) < \infty$ & $S_n = X_1 + \cdots + X_n$ then

$$P(\max_{k \le n} |S_k - E(S_k)| > x) \le \frac{\operatorname{Var}(S_n)}{r^2}$$

The next result takes many guises, but you should set the idea from examples shown.

Theorem 0.9. (Fubini-tonelli)

Suppose either that $g(x,y) \ge 0$ for all x,y or the quantities below are <u>finite</u> when g(x,y) is replaced with |g(x,y)|.

(i)
$$\iint g(x,y)dxdy = \iint g(x,y)dydx$$

.

(ii)
$$\int \sum_{n=0}^{\infty} g(n,y) dy = \sum_{n=0}^{\infty} \int g(n,y) dy$$

.

(iii)
$$\int E(g(X,y))dy = E(\int g(x,y)dy)$$

•

(iv)
$$\sum_{n=0}^{\infty} E(g(X,n)) = E(\sum_{n=0}^{\infty} g(X,n))$$

.

(v)
$$\sum_{n=0}^{\infty} \sum_{k=0}^{\infty} g(n,k) = \sum_{k=0}^{\infty} \sum_{n=0}^{\infty} g(n,k)$$

.

(vi)
$$\iint g(x,y)F(dx)G(dy) = \iint g(x,y)G(dy)F(dx)$$

, where the latter are lebesgue-stieltjes integrals. (When $g(x,y) \ge 0$ all x,y the double integrals/sums/etc above can equal ∞ , on both sides.)

We rely heavily on conditional probabilities and conditional expectation. Like expectation, however, we want to discuss & use them without being restricted to joint discrete distributions or joint continuous distributions. We provide here, without elaboration, a suitably general definition.

Definition. 0.9. Let Y be a r.v. such that E(Y) exists, and X be a random element (r.v., r. vector, stochastic process). The conditional expectation of Y given X, denoted E(y|X) is

a function of the random variable X, say h(x), (Note: this also means E(Y|X) is itself a r.v.) such that

$$E(g(x)Y) = E(g(x)E(Y|X)) \tag{*}$$

for all (measurable) bounded real-valued functions g(x). ("measurable" just means $\{x: g(x) \le y\}$ is a Borel set, which is a technical restriction not usually of much concern to us.)

Conditional Probability $P(y \in A|X) = E(\mathbb{1}_A(y)|X)$.

The definition does not tell us how to compute E(y|X); it merely says that we can define it. Note that E(y|X) is a random variable (but it must be a function of X). Often we may have a way to define or compute h(x) = E(y|X = x) suitably and then set E(y|X) = h(X).

Properties of conditional expectation

Theorem 0.10. Given X, and r.v.'s Y, Y_1, Y_2, \cdots (w/ means)

(i)
$$Y_1 \leq Y_2 \implies E(Y_1|X) \leq E(Y_2|X)$$
.

(ii)
$$E(aY_1 + bY_2|X) = aE(Y_1|X) + bE(Y_2|X)$$
.

- (iii) MCT, DCT & Fatou's lemma all hold, cond. on X.
- (iv) The formula (*) in **Theorem.0.6.** holds whenever whenever the expectations exist (not just for bounded g(x)).
- (v) (Pull-Out) E(g(x)y/X) = g(X)E(y|x), if it exists.
- (vi) (In)equalities for expectation (Jensen, etc.) all hold, cond. on X.
- (vii) If Y is independent of X then E(Y|X) = E(Y).

(viii)
$$y = h(x)$$
 then $E(y|x) = y$

(ix) (Tower)
$$E(y|x) = E(E(y|x,z)|x) = E(E(y|x)|x,z)$$
.

Basically, we can work with conditional expectations in an intuitive fashion based on familiarity with definitions for the case (x, y) is jointly discrete or jointly absolutely cont.

But now we can do so without those restrictions, including the case X is a stochastic process (or a portion of one) consisting of infinitely many rv's.

Take care to use the tower property, (ix) above, correctly. It is <u>not</u> usually true that

$$E(E(y|x)|z) = E(E(y|z)|x)$$

One other caution: E(y|x) is not originally defined since P(bx=0)=1 implies E(y|x)+b(x) also satisfies the definition. We don't usually have a problem w/ this if there are only countably many r.v.s.

1 Introduction

- definitions
- examples (simple random walk, simple branching proc.)
- probability generating functions
- some analytic results

Definition. 1.1. A <u>stochastic process</u> is a collection of random variables, $\{X_t\}_{t\in\mathbb{T}}$, defined on the same probability space. The <u>state space</u> \mathbb{X} is a set of possible values for X_t and the <u>index set</u> \mathbb{T} is the set on which X_t evolves.

(a single instance of values $\{X_t\}$ is called a sample path.)

For us, X will usually be the integers Z or the real line R (or some other subset of R but it can be R^m or something more exotic.

 \mathbb{T} will usually be the nonnegative integers \mathbb{Z}_+ or the nonnegative real numbers $\mathbb{R}_+ = [0, \infty)$, but it also can be more general.

Example. 1.1. $X_t = \#$ of people infected on day t of an epidemic

 $X_{s,t} = \text{temperature at point } s \text{ on the globe and time } t$

X(A)=# of fireant colonies in a region A (here $\mathbb T$ is a collection of sets, including all open & closed sets)

Stochastic processes have many applications:

e.g. signal process, random networks, queues, climate, earthquakes, insect populations, stock market, economics, epidemics, and so on.

Note: I generally use upper case for random variables and lower case for actual values.

Just as we do for a random variable & a random vector, we want to discuss the <u>distribution</u> of a stochastic process. But, with infinitely many rv's, we cannot simply express it as an ordinary function (such as a cdf). Instead we have

Theorem 1.1. The <u>distribution</u> of a real-valued stochastic process $\{X_t\}$ (that is, how we can calculate probabilities & expectations for it) is uniquely determined by its finite-dimensional distributions

$$F_{t_1,\dots,t_n}(x_1,\dots,x_n) = P(X_{t_1} \le x_1,\dots,X_{t_n} \le x_n)$$

The idea is that any probability about the process <u>as a whole</u> can be computed in terms of limits involving finite-dim dist's (or prob's of finitely many rv's).

A corollary of this theorem is that, in principle, it is always possible to construct a process from its distribution in terms of functions of independent r.v's.

• ex.

$$P(\sup_{n\geq 0} X_n \leq x) = \lim_{n\to\infty} P(\bigcap_{k=1}^n \{X_k \leq x\}) = \lim_{n\to\infty} F_{1,\dots,n}(x,\dots,x)$$

•

 $P(\text{the limit until } X_t \text{ first exceeds } x \text{ is finite}) = \sum_{n=0}^{\infty} P(\max_{k < n} X_k \leq x < X_n)$

(Respectively: limit of sums w/ finitely many terms, probability for finitely many r.v's)

(Note: the concept applies to all stochastic processes, whether real-valued or not)

<u>key point</u> The individual random variables in a stochastic process are <u>not independent</u>. Thus the primary problem is to model their dependence. This can be done in various ways.

- by construction from independent random variables e.g. $X_t = \sum_{j=0}^{\infty} \lambda_j \epsilon_{t-j}$, where the ϵ_t 's are iid.
- by conditional probabilities

e.g.
$$P(X_t = j | X_{t-1} = i) = {i \choose j} p^j (1-p)^{i-j}, i \ge j \ge 0$$

- by correlation structure (which may not be sufficient to characterize the process) e.g. $cov(X_t, X_{t-1}) = p^j$

Each approach offers insight into the process, so it is useful to study all three. Constructive versions of the process are also useful for simulations. Covariances are useful for statistics.

Example. 1.2. Simple Random Walk (SRW) Let S_n be a location at time n, on the integers \mathbb{Z} and suppose each new location is either up one or down one, at random, form the last location. Specifically,

$$S_0 = 0$$

$$S_{n+1} = \begin{cases} S_n - 1 & \text{w.p. } 1 - p \\ S_n + 1 & \text{w.p. } p \end{cases}, 0$$

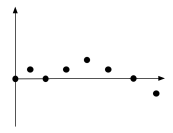


Figure 1: Simple Random Walk (SRW)

Note that, no matter what the past history of the process is, the next step depends only on the current location. We can express this by

$$P(S_{n+1} = j | S_1 = i_1, \dots, S_n = i_n) = \begin{cases} 1 - p & \text{if } j = i_n - 1 \\ p & \text{if } j = i_n + 1 \\ 0 & \text{o.w.} \end{cases}$$
$$= P(S_{n+1} = j | S_n = i_n)$$

A constructive definition is as follows: let x_1, x_2, \cdots be iid r.v's with

$$P(X_n = -1) = 1 - p$$
, $P(X_n = +1) = p$

. Then define $S_n = X_1 + \cdots + X_n$. $\{S_n\}$ is a simple random walk as defined above. (Here, $S_0 = 0$.)

Example. 1.3. Simple Branching Process (SBP)

Suppose at time 0 one individual (the prognitor) is reproductive. At time 1, this individual bears a random number $Z_{1,1}$ of descendants then becomes non-reproductive. Let $\{P_k\}_{k=0}^{\infty}$ be the probability mass function (pmf) for $Z_{1,1}$, i.e.

$$P(Z_{1,1} = k) = p_k$$

At time 2, each newly born individual randomly bears offspring, independently of each other and with the same distribution as $Z_{1,1}$. Then each becomes non-reproductive. The process continues this way, with each new individual starting its own line of descent. To represent the process constructively let $Z_{i,j}$, $i \geq 1, j \geq 1$, ne iid r.v's with pmf $\{p_k\}$, and let

 $Z_n = \text{total \# of individuals born at time } n.$

So

$$Z_0 = 1$$

$$Z_1 = Z_{1,1}$$

$$Z_2 = Z_{2,1} + Z_{2,2} + \dots + Z_{2,Z_1}$$

$$\vdots$$

$$Z_n = Z_{n,1} + \dots + Z_{n,Z_{n-1}}$$

Each Z_n is a <u>random sum</u> of Z_{n-1} id r.v's. (Define the sum to be 0 if $Z_{n-1} = 0$.) Note that each example relies on sums of independent r.v.'s. This is often the case for stochastic process (or approximately so) and at least makes a good starting point for our course. We review some basics about sums of independent r.v.'s.

Definition. 1.2. Let $\{a_n\}_{n=0}^{\infty}$ and $\{b_n\}_{n=0}^{\infty}$ be real valued sequences. Their <u>convolution</u> is the sequence (double $\{a_n\} * \{b_n\}$) with elements

$$\sum_{i=0}^{n} a_i b_{n-i} = \sum_{i=0}^{n} a_{n-i} b_i$$

.

Note that
$$\{a_n\} * \{b_n\} = \{b_n\} * \{a_n\}.$$

Also,
$$(\{a_n\} * \{b_n\}) * \{c_n\} = \{a_n\} * (\{b_n\} * \{c_n\})$$
 (check)

Convolutions have an important probabilistic interpretation, given next. Note: we write $X \sim \{a_n\}$ to indicate X has pmf $\{a_n\}$.

Lemma 1.1. Suppose X and Y are independent r.v.'s such that X has pmf $\{a_n\}$ and Y has pmf $\{b_n\}$. Then X + Y has pmf $\{a_n\} * \{b_n\}$. $(n \ge 0)$

Proof. (By asumption both X & Y are nonnegative and integer valued.)

The event $\{X+Y=n\}$ can be decomposed into disjoint events:

$${X + Y = n} = \bigcap_{i=0}^{n} {X = i, Y = n - i}$$

So

$$P(X+Y=n) = \sum_{i=0}^{n} P(X=i)P(Y=n-i)$$
 using independence
$$= \sum_{i=0}^{n} a_i b_{n-i}$$

. That is, the pmf for X + Y is $\{a_n\} * \{b_n\}$.

<u>Remark</u> If X and Y are independent and integer valued with pmf's $\{a_n\}_{n=-\infty}^{\infty}$, $\{b_n\}_{n=-\infty}^{\infty}$, rep. then

$$P(X + Y = n) = \sum_{i=-\infty}^{\infty} a_i b_{n-i}$$
 (which exists since each sequence is summable)

(A probabilistic distribution is <u>absolutely continuous</u> if it has a nonnegative density f(x) such that $\int_{-\infty}^{\infty} f(x)dx = 1$. [There are continuous distributions w/o densities.]) Or if X and Y are independent with absolutely continuous distributions having densities $f_X(x)$ and $f_Y(y)$, rep., then X + Y has density

$$f_X * f_Y(Z) = \int_{-\infty}^{\infty} f_X(x) f_Y(z - x) dx$$

. (Note: convolution formulas are context dependent, so beware!)

<u>example</u> Suppose $X \sim Poisson(\lambda), Y \sim Poisson(\mu)$. So

$$a_n = P(X = n) = \frac{\lambda^n e^{-\lambda}}{n!}, n \ge 0, b_n = P(Y = n) = \frac{\mu_n e^{-\lambda}}{n!}, n \ge 0$$

Then,

$$P(X+Y=n) = \sum_{i=0}^{n} a_i b_{n-i} = \dots = \frac{(\lambda+\mu)^n e^{-(\mu+\lambda)}}{n!}, n \ge 0, \text{ (check)}, \quad \text{and } X+Y \sim Poisson(\lambda+\mu).$$

Note: If X_1, \dots, X_n are iid with pmf $\{p_k\}$ we will denote the pmf of $S_n = X_1 + \dots + X_n$ by

$$\{P_k^{*n}\} = \{p_k\} * \cdots * \{p_k\}$$
 (n-fold convolution)

. Computing convolutions can be tedious, difficult or even impossible. Alternatively, one may use a transform. (there several different kinds of transforms.)

Definition. 1.3. (a) The generating function (gf) for a sequence $\{a_n\}_{n\geq 0}a_nS^n$ is

$$A(S) = \sum_{n=0}^{\infty} a_n S^n,$$

(defined for all S such that $\sum_{n=0}^{\infty} |a_n| S^n < \infty$.)

(b) The probability generating function (pgf) for a pmf $\{p_n\}_{n\geq 0}$ is its gf:

$$P(S) = \sum_{n=0}^{\infty} p_n S^n$$

Remark If X is a nonnegative integer valued r.v. with pmf $\{P_n\}$. Then $P(s) = E(s^x), |s| \le 1$. $(s \ge 0 \text{ suffices here})$

It often is more <u>convenient</u> & <u>illuminating</u> to use this expectation notation. examples

• $X \sim Poisson(\lambda)$

$$E(s^x) = P(s) = \sum_{k=0}^{\infty} \frac{\lambda^k e^{-\lambda} s^k}{k!} = e^{\lambda s} e^{-\lambda} = e^{-\lambda(1-s)}$$

• $X \sim binomial(n, p)$

$$E(s^{x}) = \sum_{k=0}^{n} {n \choose k} p^{k} (1-p)^{n-k} s^{k} = (ps+1-p)^{n}$$

• $X \sim geometric(p)$

$$E(s^{x}) = \sum_{n=0}^{\infty} p(1-p)^{k} s^{k} = \frac{p}{1 - (1-p)s}$$

Some useful power series to know are

$$\sum_{n=k}^{\infty} z^n = \frac{z^k}{1-z} \quad \text{if } |z| < 1$$

$$\sum_{n=0}^{m} {m \choose n} b^{m-n} z^n = (z+b)^m$$

$$\sum_{n=0}^{\infty} \frac{1}{n!} z^n = e^z$$

$$\sum_{n=0}^{\infty} \frac{(-1)^{n-1}}{n} z^n = \log(1+z) \quad \text{if } z > -1$$

An example solving a gf (rational polynomial, partial fractions)

$$A(s) = \frac{1+s+2s^2}{2-3s+s^2} = \frac{1+s+2s^2}{(2-s)(1-s)} = \frac{a+bs}{2-s} + \frac{c+ds}{1-s}$$