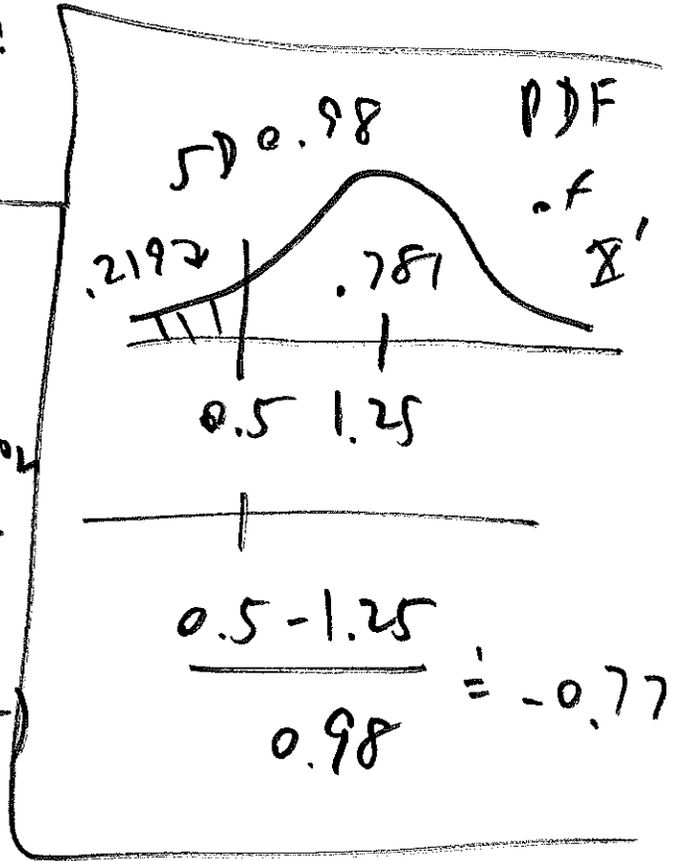


Improved approximation obtained by paying attention to the edges of the histogram ($\frac{M}{n}$) bars:

Normal approximation with continuity correction

$$\begin{aligned}
 P(X \geq 1) &\doteq 1 - P(X' < 0.5) \\
 &\doteq 1 - .219 \\
 &\doteq 0.781
 \end{aligned}$$



(correct answer 0.76; much better approx.)

Markov Chains

Recall the definition of a stochastic process:

Def. A sequence of rvs X_1, X_2, \dots (323)
is called a stochastic process with
discrete time parameter $t = 1, 2, \dots$.

X_1 is the initial state of the process;

$X_n, n \geq 1$ is the state of the process
at time $t = n$.

The simplest possible
discrete-time stochastic process is
an IID sequence of rvs (X_1, X_2, \dots) .

Suppose that there's a parameter θ
such that $(X_i | \theta) \stackrel{\text{IID}}{\sim}$ from some dist.

depending on θ . Q: Does this process
have a memory?

Example,
revisited

Machine with a dial from (324)
0 to 1, produces IID Bernoulli(θ)

trials X_i : The process (X_1, X_2, \dots)

does have a ^{for you} memory if θ is unknown

to you: the information that 17 out
of the first 20 trials were successes
helps you to predict X_{21} , because it's
reasonable to conclude from X_1, \dots, X_{20}
that θ is around $\frac{17}{20} = 0.85$, so X_{21} ~~is~~ ^{will}
probably ^{be} a success.

But the process

$\{(X_i | \theta), i=1, 2, \dots\}$ has no memory
once θ is known: information about

The first n trials is irrelevant to (325)
your prediction of X_{n+1} if you know

Q. An IID process $(X_i | \theta) \stackrel{\text{IID}}{\sim}$

is called a white-noise (stochastic)
process or a white noise time series.

Q: What's the next level of complexity
for discrete-time stochastic processes
up from white noise?

A: Allow X_{n+1}
to depend on X_n but not on X_{n-1}, X_{n-2}, \dots
(i.e., let the process have a short-term
memory, (1) time period back in the
past).

From now on, I'll suppress the dependence of the process on θ in the notation.

discrete-time

Def. A stochastic process is a (first-order) Markov chain if for $n = 1, 2, \dots$; b any real number; and for all possible sequences of states x_1, x_2, \dots

$$P(X_{n+1} \leq b \mid X_1 = x_1, \dots, X_n = x_n)$$

$$= P(X_{n+1} \leq b \mid X_n = x_n).$$

In other words, the only thing you need to know to simulate where the Markov chain is going next is where it is now.

(Can define higher-order Markov chains with memory of 2 or more time periods; we won't pursue that here.)

Def.

The set of values ~~the~~ Markov chain can take on is called its state space S , which may be finite or infinite.

(Can also have Markov chains unfolding in continuous time, e.g. X_t = stock price at time t = seconds, milliseconds, microseconds, ...; we also won't pursue that here.)

It's easy to write down

the joint P^T of a Markov chain with finite S :

Consequences

① (X_1, X_2, \dots) finite Markov chain \rightarrow

Def. A Markov chain with a finite state space is called a finite Markov chain.

$$P(X_1 = x_1, \dots, X_n = x_n) =$$

$$P(X_1 = x_1) \cdot P(X_2 = x_2 | X_1 = x_1) \cdot$$

$$P(X_3 = x_3 | X_2 = x_2) \cdot \dots$$

$$P(X_n = x_n | X_{n-1} = x_{n-1}).$$

Suppose you have a finite Markov chain with k

Def. possible states numbered $1, \dots, k$

(k integer ≥ 2) $\rightarrow \{P(X_{n+1} = j | X_n = i),$

$i, j = 1, \dots, k, n = 1, 2, \dots\}$ are called the transition distribution of the Markov chain.

If $P(X_{n+1}=j | X_n=i)$ is the same for all n , the transition distribution is said to be stationary (DS) (time-homogeneous). If

the Markov chain does have a stationary transition distribution, then the probabilities

$P_{ij} \triangleq P(X_{n+1}=j | X_n=i)$ completely characterize the Markov chain's

behavior.

in a matrix called the transition matrix.

Can arrange the P_{ij} to state P_{ij}

$$\begin{matrix}
 & \begin{matrix} 1 & 2 & \dots & k \end{matrix} \\
 \begin{matrix} P_1 \\ P_2 \\ \vdots \\ P_k \end{matrix} & = & \begin{matrix} \text{from} \\ \text{state} \end{matrix} & \begin{bmatrix} P_{11} & P_{12} & \dots & P_{1k} \\ \vdots & \vdots & \ddots & \vdots \\ P_{k1} & P_{k2} & \dots & P_{kk} \end{bmatrix}
 \end{matrix}$$

All of the elements of \underline{P} are non-negative (they're probabilities), and all of the row sums are 1 (because the chain has to go somewhere), i.e.

$$\sum_{j=1}^k p_{ij} = 1 \text{ for all } i = 1, \dots, k. \quad \text{Def.}$$

matrix versus quaternion

A square matrix $\underline{P}_{k \times k}$ with non-negative entries and ^{all} row sums equal to 1 is called a stochastic matrix.

~~(Dependent/recursive)~~

Example } Gene inheritance is Markovian.
all we need to know to predict you is the genetic story of your parents

(your grand parents, ..., are irrelevant) (33)

Suppose that

A gene of interest to you has two alleles, A and a

Then a state in

the Markov chain is of the form

{ allele 1 from parent 1, allele 2 from parent 1, allele 1 from parent 2, allele 2 from parent 2 }, for

example {Aa, Aa}.

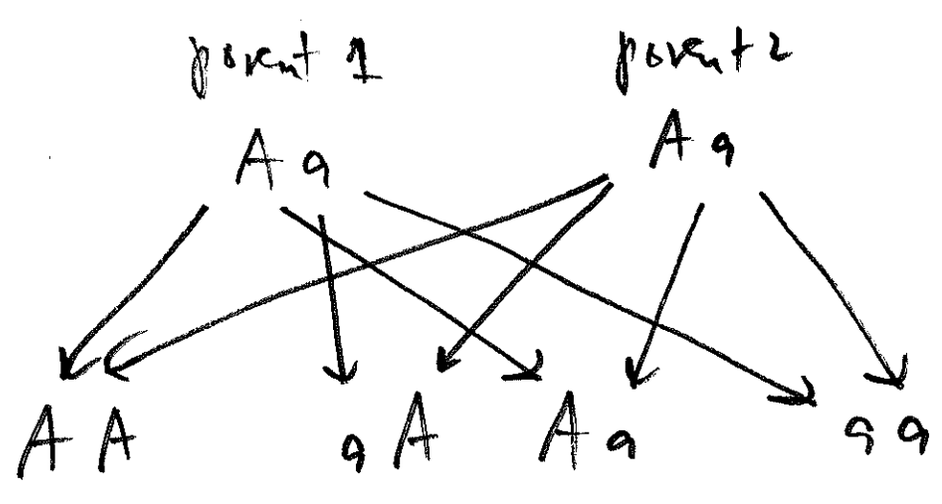
Ignoring order

(because it's irrelevant in inheritance),

there are 6 possible states: {AA, AA}

{AA, Aa}, {AA, aa}, {Aa, Aa}, {Aa, aa}

and {aa, aa}.

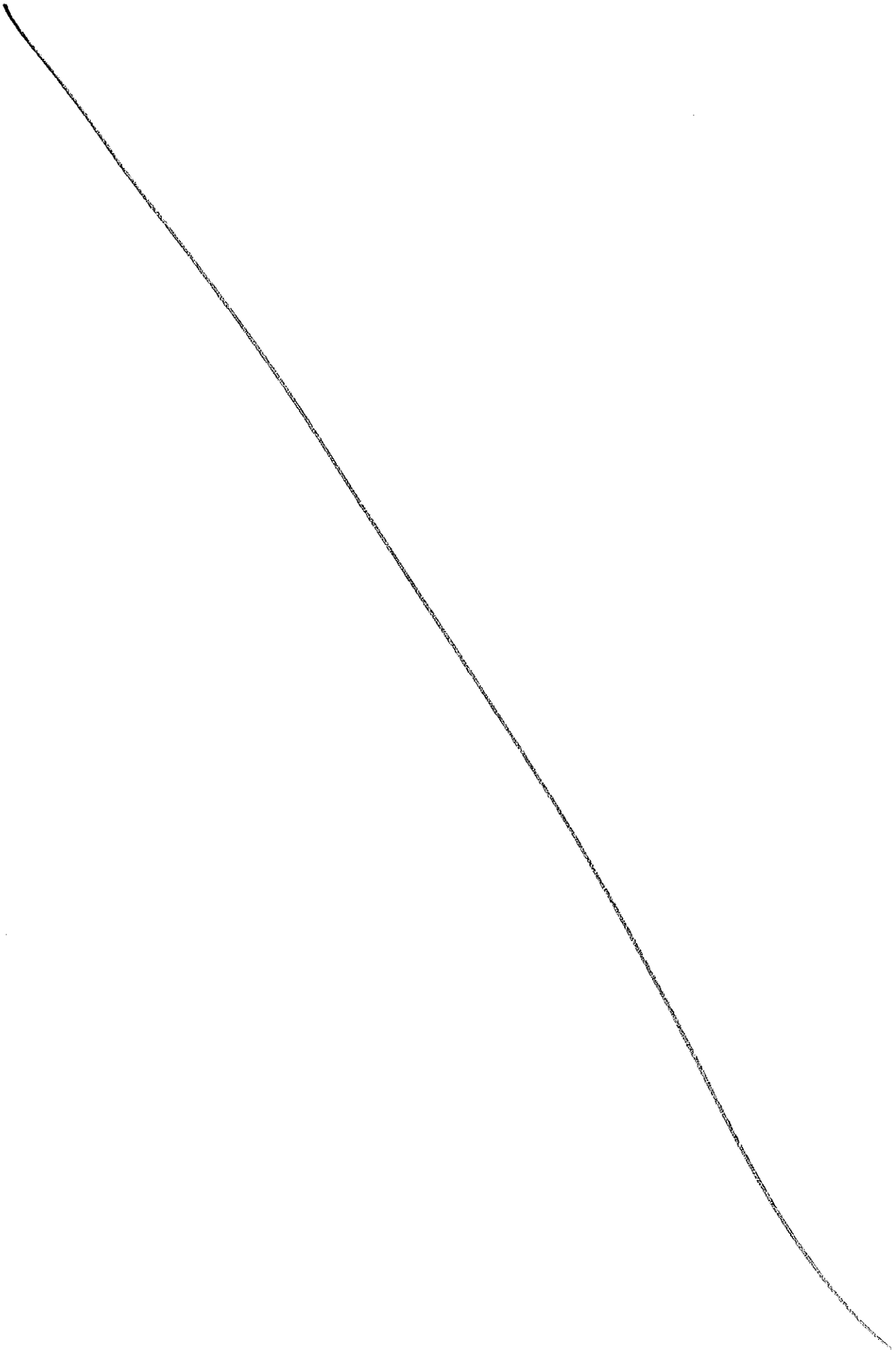


one possible inheritance sequence

offspring gets A or a from parent 1 and A or a (independently) from parent 2, each with probability $\frac{1}{2}$

Transition matrix

from \ to	{AA, AA}	{AA, Aa}	{AA, aa}	{Aa, Aa}	{Aa, aa}	{aa, aa}
{AA, AA}	1	0	0	0	0	0
{AA, Aa}	$\frac{1}{4}$	$\frac{1}{2}$	0	$\frac{1}{4}$	0	0
{AA, aa}	0	0	0	1	0	0
{Aa, Aa}	$\frac{1}{16}$	$\frac{1}{4}$	$\frac{1}{8}$	$\frac{1}{4}$	$\frac{1}{4}$	$\frac{1}{16}$
{Aa, aa}	0	0	0	$\frac{1}{4}$	$\frac{1}{2}$	$\frac{1}{4}$
{aa, aa}	0	0	0	0	0	1



Example / (random walk) You're watching 334

a particle move around on the

integers $\mathcal{S} = \{ \dots, -2, -1, 0, 1, 2, \dots \}$

over time: here are the rules:

whenever it is at time $t = n$,

it moves left 1 unit with prob p_1 ,

—— right 1 unit —— p_3 ,

and it stays where it is with prob p_2 ,

where $0 < p_i < 1$ and $\sum_{i=1}^3 p_i = 1$ This is

clearly a Markov chain (why?);

what is its transition matrix?

	to → ...	-2	-1	0	1	2	...
from ↓	⋮	⋮	⋮	⋮	⋮	⋮	⋮
-2	...	p_2	p_3	0	0	0	...
-1	...	p_1	p_2	p_3	0	0	...
0	...	0	p_1	p_2	p_3	0	...
1	...	0	0	p_1	p_2	p_3	...
2	...	0	0	0	p_1	p_2	...
⋮	⋮	⋮	⋮	⋮	⋮	⋮	⋮

= P

This is an example of a band matrix, in which the only non-zero entries are on the ^{main} diagonal and 1 diagonal either way from the main diagonal; since there are only 3 non-zero diagonals, P is said to be tridiagonal.

Moreover, all of the main diagonal entries are the same (p_2); all of the entries 1 diagonal ~~above~~ ^{below} are also the same (p_1); and all of the entries 1 diagonal above are also the same (p_3).

Such matrices are called Toeplitz

(named after Otto Toeplitz, (1881-1940) a German mathematician who was fired by the Nazis from his university position in 1935 for being Jewish. ^(died of tuberculosis at 58) Q:

Start this process, which is called a random walk, at 0 & let it go; where is the particle likely to be at time n , n large?

A: Suppose, for example, that $(p_1, p_2, p_3) = (0.1, 0.3, 0.6)$. Then you would expect the particle

(337)

to drift off to $+\infty$. Similarly,

$(p_1, p_2, p_3) = (0.5, 0.25, 0.25)$ should yield a drift to $-\infty$. $(p_1, p_2, p_3) = (\frac{1}{3}, \frac{1}{3}, \frac{1}{3})$?

Can show that as $n \rightarrow \infty$ every integer is visited infinitely many times, and the expected time you must wait for the chain to return to 0 (having started there) is also infinite.

The infinite random walk evidently has "too much freedom" to move around to get interesting results; let's bound it.

Back to a general finite Markov chain

Let $p_{ij}^{(m)} = P(\text{chain moves from } i \text{ to } j \text{ in } m \text{ steps})$ (339)

Theorem

$$= P(X_{n+m} = j \mid X_n = i)$$

Finite Markov chain with stationary transition distributions & transition

matrix $\underline{P} \rightarrow p_{ij}^{(m)}$ is just the (i, j)

entry of the matrix \underline{P}^m , which

is called the m-step transition matrix

of the Markov chain.

Genetic example, continued

$\{AA, AA\}$ has the property that once the chain is in that state, it can't

go anywhere else; so does $\{aa, aa\}$ (34)

This occurs for a state i when $p_{ii} = 1$.

Def. Any state with $p_{ii} = 1$ is

called an absorbing state.

Notice

that in this genetic Markov chain, states 2-5 all have positive probability of moving to state 1 in 2 steps, and the same is true of moving to state 6 in 2 steps.

It follows that,

if the chain is run long enough (simulating many generations), it will either end up

in state $\{AA, AA\}$ or in state $\{aa, aa\}$ (341)

Markov chains that settle down to a single ^{stable} long-run distribution are especially important in contemporary Bayesian computation; the long-run stable distribution is called the equilibrium distribution of the chain.

Important note on terminology

They call this distribution the stationary dist. of the chain, but this choice is unfortunate because they've already used stationary to mean something else:

DS: (IF $P(X_{n+1}=j | X_n=i)$ is the same for all n , DS say) that the transition distribution is stationary; other people call this time-homogeneous.

I'll use equilibrium distribution for the long-run behavior of Markov chains that settle down into a stable long-run story.

Where should the Markov chain start?

You can either initialize a Markov chain to a deterministic value, or you can start it off by making a ^{random} draw from what's called the initial distribution of the Markov chain:

Def Any vector \underline{v} of non-negative numbers (343)
that add up to 1 is called a probability
vector; any such vector whose components
specify that a Markov chain will be in
each possible state at time 1 is referred
to as the initial distribution of the chain.

So: After 1 timestep ^(iteration), the probability
dist. over the Markov chain's possible
states is \underline{v} ; after 2 iterations
the chain's dist. is $\underline{v} P$; after $(m+1)$
iterations its dist. is $\underline{v} P^m$; it would
be nice if $\underline{v} P^m$ converged to a
unique dist. as $m \rightarrow \infty$: this would

be its equilibrium distribution. (344)

Notice something interesting: if we choose \underline{v} so that $\underline{v} \underline{P} = \underline{v}$, then

$$\underline{v} \underline{P}^2 = \underline{v} \underline{P} = \underline{v}, \quad \underline{v} \underline{P}^3 = (\underline{v} \underline{P}^2) \underline{P} = \underline{v} \underline{P} = \underline{v}$$

$$= \underline{v} \underline{P} = \underline{v}; \text{ and so } \lim_{n \rightarrow \infty} \underline{v} \underline{P}^n = \underline{v}$$

Def. Markov chain with transition

matrix \underline{P} + any probability vector \underline{v}

such that $\underline{v} \underline{P} = \underline{v}$ is an equilibrium

dist. for the Markov chain

under additional

conditions on \underline{P} , such an equilibrium

dist. will be unique (we won't fully pursue that here).

How find \underline{v} so that $\underline{v} \underline{P} = \underline{v}$? (345)

In linear algebra this is an example of an eigenvalue/eigenvector problem:

Def. Given a square matrix $\underline{P}_{k \times k}$,

any vector $\underline{v}_R \in \mathbb{R}^k$ satisfying $\underline{P}_{k \times k} \underline{v}_R = \lambda_R \underline{v}_R$

is called a right eigenvector of \underline{P} with

eigenvalue λ_R , and any vector $\underline{v}_L \in \mathbb{R}^k$

satisfying $\underline{v}_L \underline{P}_{k \times k} = \lambda_L \underline{v}_L$ is called

a left eigenvector of \underline{P} with

eigenvalue λ_L .

So, given a transition matrix P for a Markov chain, an equilibrium dist. for the chain can be found by computing the left eigenvector \underline{v}_k whose eigenvalue is 1, if such a vector exists.

Most computer routines for eigenanalysis only give you right eigenvectors, but notice that if

$\underline{v}_k P = \underline{v}_k$ then $\left(\underline{v}_k P \right)^T = \underline{v}_k^T$ ← transpose

$$\underline{v}_k P = \underline{v}_k \quad \text{then} \quad \left(\underline{v}_k P \right)^T = \underline{v}_k^T$$

$\underline{v}_k^T P^T = \underline{v}_k^T$ so we can just

eigendecompose P^T instead of P .

Genetic
example,
continued

R 's routine eigen gives (347)
the following results: P^T has

two eigenvectors whose
eigenvalues are 1: $\begin{pmatrix} 1 \\ 0 \\ 0 \\ 0 \\ 0 \\ 0 \end{pmatrix}$ and $\begin{pmatrix} 0 \\ 0 \\ 0 \\ 0 \\ 0 \\ 1 \end{pmatrix}$,
corresponding to the
two absorbing states.

This suggests that there's an entire family
of equilibrium distributions, of the
form $(p, 0, 0, 0, 0, 1-p)^T$ for
 $0 \leq p \leq 1$; and Maple verifies this conjecture.

So the earlier guess is also correct:
after many generations either one of
 $\{AA, AA\}$ or $\{aa, aa\}$ will be absorbing.

There is a special case in which a unique stationary distribution exists.

Theorem If you can find a positive integer $m \geq 1$ such that every element of P^m is strictly positive, then $\lim_{n \rightarrow \infty} P^n$ is a matrix with all rows equal to the unique stationary dist \underline{v} , and no matter what the chain's initial distribution is, its distribution after n steps converges to \underline{v} as $n \rightarrow \infty$.